# Early initiation of short-term emollient use for the prevention of atopic dermatitis in high risk infants – the STOP AD randomised controlled trial

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# Abstract

**Background** Protecting the skin barrier in early infancy may prevent atopic dermatitis (AD). We investigated if daily emollient use from birth to 2 months reduced AD incidence in high risk infants at 12 months. **Methods** This was a single-center, two-armed, investigator-blinded, randomized controlled clinical trial (NCT03871998). Term infants identified as high risk for AD (parental history of AD, asthma or allergic rhinitis) were recruited within 4 days of birth and randomised 1:1 to either twice-daily emollient application for the first 8 weeks of life (intervention group), using an emollient specifically formulated for very dry, AD-prone skin, or to standard routine skin care (control group). The primary outcome was cumulative AD incidence at 12 months. AD <6 months was diagnosed based on clinical presence of AD. The UK Working Party Diagnostic Criteria were applied when diagnosing AD between 6 and 12 months. **Results** 321 infants were randomised (161 intervention and 160 control), with 61 withdrawals (41 intervention, 20 control). The cumulative incidence of AD at 12 months was 32.8% in the intervention group vs. 46.4% in the control group, p = 0.036 [Relative risk (95%CI): 0.707 (0.516, 0.965)]. One infant in the intervention group was withdrawn from the study following development of a rash that had a potential relationship with the emollient. There was no significant difference in the incidence of skin infections between the intervention and control groups during the intervention period (5.0% vs. 5.7%, P>0.05). **Conclusions** This study has demonstrated that early initiation of daily specialized emollient use until 2 months reduces the incidence of AD in the first year of life in high-risk infants.

# Introduction

Atopic dermatitis (AD), also known as eczema, is a chronic inflammatory skin condition, characterized by dry, red and itchy skin<sup>1,2</sup>. AD usually begins in infancy and affects up to one fifth of children <sup>3,4</sup>. The pathogenesis includes impaired skin barrier function as a significant pathomechanism, along with cutaneous immune dysregulation and microbial disturbances<sup>5</sup>. Supporting this is the consistent evidence that loss-of-function mutations in the filaggrin gene (*FLG*), resulting in measurable skin barrier defects, plays a central role in the inherited risk of AD <sup>6</sup>.

Daily emollient use is a cornerstone of AD management<sup>7</sup>. Recently, the spotlight has been on the potential role of emollients in infancy in preventing AD. Prompting this were findings from two small randomised control trials (RCTs) reporting that daily emollient application from birth until 6-8 months reduced AD risk by up to 50% <sup>8,9</sup>. Unexpectedly, these findings were not replicated in two much larger scale studies  $^{10,11}$ . The Barrier Enhancement for Eczema Prevention (BEEP) trial recruited 1394 high-risk infants and randomised them to either daily emollient application for the first year or to standard skin care advice

alone<sup>10</sup>. No evidence of a protective effect of emollient use against AD at 1 or 2 years was found. The Preventing Atopic Dermatitis and Allergies in Childhood (PreventADALL) study involved baths for 5-10 minutes with added emulsified oil and cream applied to the face after the bath on at least 4 days per week from 2 weeks to 8 months and reported no effect on AD prevalence when assessed 4 months later at 12 months. Another RCT where emollient was applied daily to the face only from 0-6 months also reported no effect <sup>12</sup>. The data from these three RCTs largely contributed to the conclusion of a recent meta-analysis that skin care interventions probably do not influence AD development <sup>13</sup>. This meta-analysis used an individual participant data approach, excluding studies only providing aggregate data. In contrast, another meta-analysis including more studies found a beneficial effect of emollients in high-risk infants [RR (95% CI: 0.59 (0.43, 0.81)], but only when used up to the point of AD assessment and not when there was an interval between the treatment and the assessment <sup>14</sup>.

The BEEP and PreventADALL studies used petroleum and paraffin-based emollient formulations, <sup>10,11</sup> and while Dissanayake et al. used a more complex ceramide-based emollient, the latter study's intervention involved application to the face only <sup>12</sup>. Data from a small pilot study suggest that emollients with ingredients specifically designed to repair the skin barrier warrant further investigation <sup>15</sup>. Interventions in BEEP and PreventADALL began at a median age of 11 days and from 2 weeks and continued for 12 and 8 months, respectively <sup>10,11</sup>. Daily emollient application for an extended period in infancy places considerable additional demands on new parents and may not be feasible at a population level, especially if specialized and more expensive emollients are advised. This may be reflected in the low adherence of 27% to the intervention in PreventADALL <sup>11</sup>. We have shown that trans-epidermal water loss (TEWL) increased from birth to 2 months but stabilised thereafter <sup>16</sup> suggesting a shorter intervention period, beginning as soon as possible after birth may represent a more feasible intervention, while targeting a critical period of skin maturation.

This study aimed to investigate if daily emollient use from birth to 2 months can reduce the incidence of AD in high risk infants.

## Methodology

# Study design

Short-term Topical Application to Prevent Atopic Dermatitis (STOP AD) was a single-centre, two-armed randomized control trial that postnatally recruited newborn infants at high-risk of AD. Recruitment took place between April 2019 and November 2020 in Cork University Maternity Hospital (CUMH). Parents gave written informed consent prior to participation. Term infants were identified as high-risk if they had at least one parent with a history of AD, asthma or allergic rhinitis. Exclusion criteria were: pre-term infant (born <37 weeks), admission to the neonatal unit for issues other than feeding, receipt of antibiotics in the maternity hospital, phototherapy, sibling already recruited, other serious health conditions, severe widespread skin condition or any condition that would make the emollient use inadvisable or not possible (e.g. ankle talipes or hip dysplasia). The study was conducted in accordance with the Helsinki Declaration and was approved by the Clinical Research Ethics Committee of the Cork Teaching Hospitals [ref ECM 5 (2) 18/12/18].

#### Randomisation

Infants were randomised (1:1 allocation ratio) within 4 days of birth to either twice-daily emollient application for the first 8 weeks of life (intervention groupintervention - IG) or to standard routine skin care advice (control group - CG). Parents and research nurses and staff responsible for recruitment, taking measurements and administering questionnaires were not blinded to study allocation. The study doctor performing AD and food allergy assessments was not involved in recruitment and was blinded throughout the study. Parents in the IG were instructed to apply the emollient AVEENO® Dermexa Fast & Long Lasting Balm (Johnson & Johnson Santé Beauté France, JJSBF) twice-daily to the whole body (excluding scalp) for the first 8 weeks. This product, with oat ingredient, fatty acids and ceramides, developed specifically for very dry itchy skin AD-prone skin, was supplied free to the study by the manufacturer but is publicly available for sale and for use in this age group. Emollient use after the intervention period was at parental discretion. The CG were advised to follow the standard skin care advice given at CUMH, which does not include regular emollient use, unless indicated. Both groups were provided with AVEENO® Baby Daily Care Baby Gentle Wash (JJSBF) to be used at their discretion. Adherence was assessed using questionnaires at 2, 4 and 8-week and diaries completed over the intervention period. Adherence to the intervention was defined as using an emollient at least once daily. Contamination in the CG was defined as emollient on four or more days per week.

#### Study visits and procedures

Study visits were at baseline (pre-discharge, within 4 days of birth) and at approximately 2, 4 and 8 weeks and at 6 and 12 months. These involved questionnaires on feeding, health, skin care and bathing; repeat measurements of weight; TEWL and natural moisturising factor (NMF) and monitoring of skin health.

## Skin barrier assessments

TEWL was measured on the volar forearm using a closed chamber system vapometer (Delfin Technologies, UK) after acclimatization to room conditions for at least ten minutes.

# Filaggrin genotyping/stratification

As a proxy for FLG genotyping, NMF was measured *in vivo*, non-invasively, at a depth of 25 µm in the stratum corneum of the thenar eminence by Near-Infrared Raman spectroscopy (NMF-scan, RiverD International B.V., Rotterdam, The Netherlands), a method that has previously been shown to be an excellent proxy for FLG genotyping<sup>17–19</sup>. Formal FLG genetic testing as performed as follows: buccal swabs were collected using Isohelix SK-3S swabs and BFX/S1/05/50 buccal fix tubes (Cell Projects ltd). Filaggrin genotyping was performed at the A\*STAR Skin Research Labs using Microfluidics PCR for full coverage of FLG repeat alleles using a method described previously<sup>20</sup>.

## Atopic dermatitis assessment

Parents were routinely encouraged to report skin concerns to the study team. Suspected cases of AD were reviewed by the blinded investigator at the earliest opportunity. Cases of AD <6 months were diagnosed based on the presence of AD, assessed either in person or via photographs when an in-person assessment was not possible. The UK Working Party Diagnostic Criteria (UKWPDC) were applied when diagnosing AD between 6 and 12 months <sup>21</sup>. AD extent and severity were evaluated (blinded) using SCORing Atopic Dermatitis (SCORAD) for infants [?]6 months <sup>22</sup>. During the high-level COVID-19 restrictions, AD assessments, including SCORADs, were completed remotely by the blinded investigator using photographs and video links (see supplementary data for impact of COVID-19 on study). AD cases were treated with a standardized treatment program which included advice on emollient use and a topical steroid treatment, where required.

#### Food allergy assessment

Parents were advised to introduce common food allergens including egg, dairy and peanut early during weaning, as per national guidelines. Suspected cases of food allergy were clinically assessed by the blinded investigator and skin prick testing (SPT) was performed, as indicated. Infants also had SPT to egg, dairy and peanut if they hadn't safely consumed these foods by 12 months. Where deemed necessary by the allergy team, those with a positive SPT or a reaction suggestive of food allergy were invited for an oral food challenge (OFC).

# Outcomes

The primary outcome was cumulative incidence of AD at 12 months. Secondary outcomes included AD incidence at 6 months, cumulative incidence of sensitization to food at 12 months and the evolution of TEWL and NMF between 0-12 months.

# Sample size

The target sample size was 242 (n = 121 per group). This would provide 80% power at a 95% confidence level to detect a 50% reduction in cumulative AD at 12 months from 30% to 15%. The expected AD rate of 30% in this high risk group was based on data from a previous Irish birth cohort study in the same geographical area <sup>23</sup>.

#### Data analysis

Data were analysed based on a "as randomized, complete-case" approach, where those missing an AD outcome were excluded. Groups were analysed for the primary outcome as randomised, regardless of adherence to study allocation. Sensitivity per-protocol analyses were conducted based on adherence data from questionnaires and diaries. Using parent-reported emollient data from questionnaires at 2, 4 and 8 weeks, participants in the IG were included in the per-protocol analysis if they reported at least once-daily emollient use at each of the three time-points. CG participants were included if they reported emollient use of <4 days per week at each time-point. In the diary per-protocol analysis, participants in the IG were included if they recorded emollient use on [?]90% of days in the 8-week recording period, equating to >6 days a week. CG participants were included if they used emollient on [?]43% of the days (<4 days a week). Additional sensitivity analyses were conducted including those in the IG who reported emollient use on [?]4 days a week and those who in the CG who used emollient on <4 days3 days a week to more closely align with the adherence definition used in the BEEP study<sup>10</sup>.

#### Results

## **Recruitment**/retention

A total of 3059 infants were screened for eligibility between April 2019 and November 2020, of whom 321 were randomised (161 to intervention and 160 to control), Figure 1. Baseline characteristics were balanced across the groups (Table 1). There were 61 withdrawals (41 intervention and 20 control, 19% attrition), with the majority (80%) occurring before the 2 week visit. The mean (SD) age at randomization was 1.9 (0.9) days.

## **Protocol adherence**

In the questionnaires, most parents in the IG reported applying emollient at least once daily in the first 8 weeks; 2 weeks: 89%, 4 weeks: 91.7%, 8 weeks: 86.6% (Table 2). Twice-daily application was reported by 63.3% at 2 weeks, 69.2% at 4 weeks and 73.1% at 8 weeks. Of those in the IG with questionnaires at all three time-points (n = 114), 89 (78.1%) reported daily emollient use at all three time-points. Less than 20% of the CG reported emollient use on [?]4 days per week at any of the time-points; 19% at 2 weeks, 17.5% at 4 weeks and 13.5% at 8 weeks. Of those in the CG with questionnaires at all time-points (n = 132), 90 (68.2%) reported using emollients on < 4 days per week at all three time-points. There was no significant difference in bathing frequency between the groups over the intervention period (See supplementary data and Supplementary Table 7).

Diaries measuring adherence were returned by 95% (114/120) of the IG and 82.1% (115/140) of the CG. The mean (SD) age that emollient use started in the IG was 3.5 (1.5) days and 41.2% (47/114) reported that they applied emollient at least once on [?]90% of recording days (>6 days/week). A further 41.2% (47/114) reported emollient use for [?]75% of recording days (>5 days/week). Eighty percent (92/115) of the CG applied an emollient on [?]43% of the recording days ([?]3 days a week).

There was no significant difference in the prevalence of regular emollient use ([?]4 days a week) between the groups at 6 and 12 months (intervention vs. control: 29.6% vs 29.7%, p = 1.000 at 6 months and 28.4% vs. 25.8%, p = 0.868 at 12 months) (See Supplementary data and Supplementary Table 8).

#### Safety

No family sought emergency medical assessment related to the study intervention. Parent-reported skin infections during the 8-week intervention period occurred in 5% (6/120) of the IG and 5.7% (8/140) of the CG. One IG infant was advised to stop applying the emollient after developing a rash that had a potential

temporal relationship with the emollient and was withdrawn from the study. Two suspected reactions to the study emollient were investigated and confirmed as having no relationship.

#### **Primary outcome**

The cumulative incidence of AD at 12 months was 32.8% in the IG vs. 46.4% in the CG, p = 0.036 [Relative risk (RR) (95% CI): 0.707 (0.516, 0.965), Figure 2]. The point prevalence of AD at 12 months, where the child met the UKWPDC at the assessment, was 20.5% in the IG vs. 38.2% in the IG, p = 0.003 [RR (95% CI): 0.536 (0.354, 0.813)].

#### Secondary AD outcome

The cumulative incidence of AD at 6 months was 18.3% in the IG vs. 36.4% in the CG, p = 0.002 [RR (95%CI): 0.503 (0.325, 0.779)]. The point prevalence at 6 months was 18.3% in the IG and 35.0% in the CG, p = 0.004 [RR (95%CI): 0.524 (0.337, 0813)], Table 3.

Time-to-event survival analysis using the Kaplan- Meier method demonstrates that the IG maintained ADfree skin for a longer period in the first 12 months than the CG (p = 0.016, log-rank test, Figure 3). Of those with AD outcome data at 6 and 12 months (n = 117 intervention, n = 137 control), 7.7% of IG and 8.0% of CG infants were diagnosed at [?]6 months, but no longer met the criteria at 12 months (p = 1.0). The prevalence of AD onset between 6 and 12 months was 13.7% and 9.5% in the IG and CG, respectively (p = 0.397) and 10.3% vs. 29.2% met the criteria at both 6 and 12 months (p < 0.001), (See Supplementary Figure 1). SCORADs were completed for those [?]6 months at diagnosis (n = 55). There was no significant difference in SCORAD total scores at diagnosis between the groups [median (IQR) SCORAD: IG 11.3 (8.0, 18.4), vs. CG 12.3 (7.4, 16.0), p = 0.888].

A similar, but non-significant relative risk was observed for the primary outcome in the per-protocol analyses [Questionnaire per-protocol analysis RR (95%) CI: 0.713 (0.501, 1.014), P = 0.078); Diary per-protocol analysis RR (95% CI): 0.745 (0.474, 1.173), P = 0.253], (See Supplementary Tables 1-4).

### Food allergen sensitization

All infants had been introduced to dairy and almost all had been introduced to egg (99.6%) and peanut (98.0%) by 12 months. Nine infants had a positive SPT to at least one food [intervention; 3.3% (4/120), control; 3.6% (5/120), p =1.0].

## **TEWL and NMF evolution**

There were no significant differences in TEWL or Thenar NMF between the intervention and control groups at birth, 2, 4, 8 weeks or at 6 and 12 months (See Supplementary Tables 5 & 6).

#### Discussion

In this RCT in high-risk infants, we found that daily emollient use initiated in the first week of life until 2 months is associated with a significant reduction in the cumulative incidence of AD at 12 months. Daily emollient use was associated with a 50% and 29% reduction in the risk of the cumulative incidence of AD at 6 and 12 months, respectively. Similar risk reductions were observed in the per-protocol analyses where only those in the intervention and control groups were included if they used emollients at least once daily and <4 days a week, respectively. However, these were not significant for the primary outcome which may be due to the conservative adherence criteria applied and thus, lower numbers included in the analysis and therefore lower power to detect differences between the groups.

While some AD cases diagnosed before 6 months had resolved by 12 months, there was no difference in transient cases between the groups. As we did not collect longer term data, we cannot exclude the possibility that the intervention may have only delayed the onset of AD beyond 12 months. A recent meta-analysis reported a protective effect of emollients but only when there was no interval between the emollient treatment and AD assessment <sup>14</sup>. However, there was significant heterogeneity between the four studies included in

that analysis. In our study, a 29% reduction in the risk of cumulative AD at 12 months, was maintained 10 months after the intervention.

Our findings are at variance with recent findings from two large RCTs, where no evidence of a protective effect of emollient use in the first year against AD was found <sup>10,11</sup>. Among the most notable differences between these RCTs and ours was the timing of the intervention. The treatment in STOP AD began within days of birth during a dynamic period of skin maturation and adaption to the dramatic environmental changes of life ex utero. In STOP AD, infants were randomised within 4 days of birth with the IG advised to begin the emollient treatment immediately. In BEEP, the median (IQR) age that emollient use began was 11 days (7, 17) days, with only 89% starting emollient application before 3 weeks. In PreventADALL, the intervention began from 2 weeks of age.

The emollients used in BEEP and PreventADALL were basic petroleum and paraffin-based formulations, respectively. The emollient used in this study consists of a formulation with added ceramides developed specifically for very dry itchy skin. Two small studies that also used more complex ceramide-rich emollients reported non-significant trends towards a protective effect against AD <sup>15,24</sup>. Following one of these <sup>15</sup>, a larger scale RCT, the PEBBLES study, involving twice-daily application of the same ceramide-based emollient from 0-6 months is ongoing<sup>25</sup>. Here we showed a reduced risk of AD at 12 months with a short 2-month intervention period, which may represent a more feasible and family friendly strategy for AD prevention.

Our high adherence rates demonstrate the feasibility of implementing a regimen of daily emollient use during the first 2 months of life. Adherence rates using the diaries were lower than reported on the questionnaires but 82.4% still reported using emollients on [?]75% of days equating to over 5 days a week. While infants in this study were followed closely during the intervention period, similar rates of adherence were observed in BEEP which involved limited contact, but used a less strict definition for adherence (emollient use [?]3 days/week)<sup>10</sup>. Only 27% of the IG fully adhered to the protocol in PreventADALL which may have influenced the absence of a protective effect <sup>11</sup>.

While we did assess food allergy outcomes, this study was not powered to detect a reduction in food allergy risk. Unlike BEEP, where a non-significant increase in food allergy in the IG has been prominently reported (15), we found no difference in the prevalence of food allergy between the groups. While we did not use SPTs to screen for food allergy, almost all infants had tried the most common food allergens - milk, egg and peanut - by 12 months, so the rate of food sensitization and allergy reported is likely reflective of the true rate in our groups. BEEP reported a higher rate of skin infections in the IG, with suggestions of the possibility of greater pathogen exposure with emollient application <sup>10</sup>. We did not find evidence of an increased risk of skin infections with short-term emollient use.

Despite the reduction of AD risk in the IG, there was no difference in TEWL throughout the first year between the groups. Other studies on emollient use during infancy reported a similar absence of an effect of the intervention on TEWL <sup>15,24</sup>. TEWL measurements are influenced by environmental factors and more crucially for infants, subject-specific parameters including stress and crying<sup>26</sup>. This may have affected our ability to detect differences between the groups.

The major strength of this study is the initiation of emollient use within days of birth in the IG. Other strengths include the close follow-up of infants, a high rate of adherence in the IG and a low rate of contamination in the CG.

A limitation to this study is that in response to the COVID-19 pandemic, many AD diagnoses were made remotely. To mitigate this, detailed information and photographs were collected when making a diagnosis. SCORAD assessments were also completed remotely, which may have affected assessments of AD severity. Validated diagnostic criteria could not be applied when diagnosing earlier onset AD (<6 months), where cases were diagnosed based on presence of AD lesions. However, of the 73 infants diagnosed with AD [?]6 months, 71 (97.3%) met the UKWPDC at 6 months. The prevalence of cumulative AD in this group was higher than expected based on the rates among infants with parental history of atopy in an Irish birth cohort <sup>23</sup>. A possible explanation for this is the *a priori* recruitment of high-risk infants and the close monitoring of skin health in this study. Only a third (32.1%) of those eligible for this study were recruited. One of the main reasons for the refusal to participate was the demanding follow-up schedule involved, particularly during the intervention period that started before going home with their newborn baby, suggesting that more motivated individuals were recruited. We also had a higher rate of withdrawals in the first two weeks of life, particularly in the intervention group, mainly due to withdrawal of consent and not due to early onset of AD by this time. This is a consideration in assessing the feasibility of advising daily emollient use in the early postnatal period to a more general population.

We have demonstrated that early initiation of daily specialized emollient use until 2 months reduces the incidence of AD in the first year of life in high-risk infants. The mechanisms behind this are unclear but analysis of microbiome diversity and inflammatory biomarkers in a subgroup of this study is ongoing and may provide further information. While several recent studies do not support a protective effect of emollient use in infancy, future studies should examine the use of more complex emollients directed at enhancing the skin barrier, while identifying a treatment window that is both effective and acceptable to parents.

# References

1. Weidinger S, Beck LA, Bieber T, Kabashima K, Irvine AD. Atopic dermatitis. *Nat Rev Dis Primer*. 2018;4(1):1. doi:10.1038/s41572-018-0001-z

2. Langan SM, Irvine AD, Weidinger S. Atopic dermatitis. Lancet Lond Engl . 2020;396(10247):345-360. doi:10.1016/S0140-6736(20)31286-1

3. Odhiambo JA, Williams HC, Clayton TO, Robertson CF, Asher MI. Global variations in prevalence of eczema symptoms in children from ISAAC Phase Three. J Allergy Clin Immunol . 2009;124(6):1251-1258.e23. doi:10.1016/j.jaci.2009.10.009

4. Nutten S. Atopic dermatitis: global epidemiology and risk factors. Ann Nutr Metab . 2015;66 Suppl 1:8-16. doi:10.1159/000370220

5. Leung DYM. New insights into atopic dermatitis: role of skin barrier and immune dysregulation. Allergol Int Off J Jpn Soc Allergol . 2013;62(2):151-161. doi:10.2332/allergolint.13-RAI-0564

6. Drislane C, Irvine AD. The role of filaggrin in atopic dermatitis and allergic disease. Ann Allergy Asthma Immunol Off Publ Am Coll Allergy Asthma Immunol . 2020;124(1):36-43. doi:10.1016/j.anai.2019.10.008

7. McAleer MA, Flohr C, Irvine AD. Management of difficult and severe eczema in childhood. BMJ . 2012;345:e4770. doi:10.1136/bmj.e4770

8. Simpson EL, Chalmers JR, Hanifin JM, et al. Emollient enhancement of the skin barrier from birth offers effective atopic dermatitis prevention. *J Allergy Clin Immunol* . 2014;134(4):818-823. doi:10.1016/j.jaci.2014.08.005

9. Horimukai K, Morita K, Narita M, et al. Application of moisturizer to neonates prevents development of atopic dermatitis. J Allergy Clin Immunol . 2014;134(4):824-830.e6. doi:10.1016/j.jaci.2014.07.060

10. Chalmers JR, Haines RH, Bradshaw LE, et al. Daily emollient during infancy for prevention of eczema: the BEEP randomised controlled trial.*Lancet Lond Engl*. 2020;395(10228):962-972. doi:10.1016/S0140-6736(19)32984-8

11. Skjerven HO, Rehbinder EM, Vettukattil R, et al. Skin emollient and early complementary feeding to prevent infant atopic dermatitis (PreventADALL): a factorial, multicentre, cluster-randomised trial. *Lancet Lond Engl*. 2020;395(10228):951-961. doi:10.1016/S0140-6736(19)32983-6

12. Dissanayake E, Tani Y, Nagai K, et al. Skin Care and Synbiotics for Prevention of Atopic Dermatitis or Food Allergy in Newborn Infants: A 2 × 2 Factorial, Randomized, Non-Treatment Controlled Trial. Int Arch Allergy Immunol . 2019;180(3):202-211. doi:10.1159/000501636

13. Kelleher MM, Cro S, Van Vogt E, et al. Skincare interventions in infants for preventing eczema and food allergy: A cochrane systematic review and individual participant data meta-analysis. *Clin Exp Allergy J Br Soc Allergy Clin Immunol*. 2021;51(3):402-418. doi:10.1111/cea.13847

14. Zhong Y, Samuel M, van Bever H, Tham EH. Emollients in infancy to prevent atopic dermatitis: A systematic review and meta-analysis. *Allergy*. Published online September 30, 2021. doi:10.1111/all.15116

15. Lowe AJ, Su JC, Allen KJ, et al. A randomized trial of a barrier lipid replacement strategy for the prevention of atopic dermatitis and allergic sensitization: the PEBBLES pilot study. Br J Dermatol . 2018;178(1):e19-e21. doi:10.1111/bjd.15747

16. Kelleher M, Dunn AG, Irvine A, Smith H, Hourihane OJ. O08 - Increased early life Transepidermal Water Loss (TEWL) values can predate atopic dermatitis in asymptomatic infants: results from the BASELINE study. *Clin Transl Allergy* . 2014;4(1):O8. doi:10.1186/2045-7022-4-S1-O8

17. Caspers PJ, Lucassen GW, Carter EA, Bruining HA, Puppels GJ. In vivo confocal Raman microspectroscopy of the skin: noninvasive determination of molecular concentration profiles. *J Invest Dermatol* . 2001;116(3):434-442. doi:10.1046/j.1523-1747.2001.01258.x

18. Ní Chaoimh C, Nico C, Puppels GJ, et al. In vivo Raman spectroscopy discriminates between FLG loss-of-function carriers vs wild-type in day 1-4 neonates. Ann Allergy Asthma Immunol Off Publ Am Coll Allergy Asthma Immunol . 2020;124(5):500-504. doi:10.1016/j.anai.2020.01.022

19. O'Regan GM, Kemperman PMJH, Sandilands A, et al. Raman profiles of the stratum corneum define 3 filaggrin genotype-determined atopic dermatitis endophenotypes. *J Allergy Clin Immunol* . 2010;126(3):574-580.e1. doi:10.1016/j.jaci.2010.04.038

20. Wong XFCC, Denil SLIJ, Foo JN, et al. Array-based sequencing of filaggrin gene for comprehensive detection of disease-associated variants. *J Allergy Clin Immunol* . 2018;141(2):814-816. doi:10.1016/j.jaci.2017.10.001

21. Williams HC, Burney PG, Pembroke AC, Hay RJ. The U.K. Working Party's Diagnostic Criteria for Atopic Dermatitis. III. Independent hospital validation. *Br J Dermatol* . 1994;131(3):406-416. doi:10.1111/j.1365-2133.1994.tb08532.x

22. Kunz B, Oranje AP, Labrèze L, Stalder JF, Ring J, Taïeb A. Clinical validation and guidelines for the SCORAD index: consensus report of the European Task Force on Atopic Dermatitis. *Dermatol Basel Switz* . 1997;195(1):10-19. doi:10.1159/000245677

23. O'Donovan SM, Murray DM, Hourihane JO, Kenny LC, Irvine AD, Kiely M. Cohort profile: The Cork BASELINE Birth Cohort Study: Babies after SCOPE: Evaluating the Longitudinal Impact on Neurological and Nutritional Endpoints. *Int J Epidemiol* . 2015;44(3):764-775. doi:10.1093/ije/dyu157

24. McClanahan D, Wong A, Kezic S, et al. A randomized controlled trial of an emollient with ceramide and filaggrin-associated amino acids for the primary prevention of atopic dermatitis in high-risk infants. *J Eur Acad Dermatol Venereol JEADV* . 2019;33(11):2087-2094. doi:10.1111/jdv.15786

25. Lowe A, Su J, Tang M, et al. PEBBLES study protocol: a randomised controlled trial to prevent atopic dermatitis, food allergy and sensitisation in infants with a family history of allergic disease using a skin barrier improvement strategy. *BMJ Open*. 2019;9(3):e024594. doi:10.1136/bmjopen-2018-024594

26. Stamatas GN, Nikolovski J, Mack MC, Kollias N. Infant skin physiology and development during the first years of life: a review of recent findings based on in vivo studies. *Int J Cosmet Sci* . 2011;33(1):17-24. doi:10.1111/j.1468-2494.2010.00611.x

 Table 1 Baseline characteristics

	Intervention $(n = 161)$	Control $(n = 160)$
	Intervention $(n = 161)$	Control $(n = 160)$
Maternal characteristics		
Age [mean (SD) years]	33.3(4.4)	34.1(4.8)
Country of birth (Ireland)	142 (88.2)	139 (86.9)
Ethnicity (white)	156(96.9)	158(98.8)
Paternal characteristics		
Age [mean (SD) years]	35.3(5.7)	35.9(5.5)
Country of birth (Ireland)	142(88.2)	143(89.4)
Ethnicity (white)	157 (97.5)	159(99.4)
Infant characteristics		
Sex (male)	79 (49.1)	85 (53.1)
Gestational age [mean (SD)	39.7(1.1)	395(11)
weeks]	00.1 (1.1)	00.0 (1.1)
Birth weight [mean (SD) kg]	36(04)	36(05)
Mode of delivery	0.0 (0.1)	0.0 (0.0)
Vaginal	98 (61.3)	103 (64 4)
Caesarean section	62 (38.8)	57 (35 6)
Age randomised [mean (SD)	19(10)	1.8(0.8)
davs]	1.0 (1.0)	1.0 (0.0)
Baseline TEWL [median (IOB)	9 31 (7 25 12 41)	9 25 (7 44 13 31)
water/m2/h]	5.51 (1.26, 12.11)	0.20 (1.11, 10.01)
Baseline NMF [median (IOB)	0.32(0.22, 0.42)	0.33(0.25, 0.41)
g/g protein]	0.02 (0.22, 0.12)	0.00 (0.20, 0.11)
Family history of atopy		
Maternal atony		
Allergic rhinitis	81 (50.3)	63 (39.4)
Atopic dermatitis	40(24.8)	56(35.1)
Asthma	54 (335)	63 (39.4)
Any maternal atony	112(69.6)	107 (67 3)
Paternal allergy	112 (05.0)	101 (01.5)
Allergic rhinitis	66 (41.8)	75(472)
Atopic dermatitis	(41.0)	45 (28.3)
Asthma	56(25.4)	53(20.0)
Any paternal atony	101(63.9)	107 (67 3)
Two parents with atopic history	52(32.0)	54 (34 2)
Participant with at least one	52(52.5) 87(540)	04(59.2)
sibling	87 (34.0)	94 (38.8)
Of which at least one sibling		
with		
Allorgia rhinitia	20(230)	91 (99 3)
Atopia dormatitia	20(23.0) 25 (40.2)	21 (22.3) 41 (42.6)
Asthma	10 (10.2)	(43.0) 12 (12 2)
Astillia FLC constraine	12 (13.0)	10 (10.0)
FLC wildtyping	06/117(99.1)	119/196 (09.1)
FLG wildtype	90/111(02.1) 91/117(17.0)	110/100 (00.1) 20/126 (16.0)
FLG null mutation (one)	21/117(17.9)	$\frac{22}{130} (10.2)$ $\frac{1}{126} (0.7)$
FLG null mutation (two)	U	1/130(0.7)

Data are n (%) unless stated otherwise.

Abbreviations : TEWL = Transepidermal water loss, NMF = Natural mositurising factor, FLG = gene encoding filaggrin

	Intervention $(n = 100)$	Intervention $(n = 100)$	Intervention $(n = 100)$	Control $(n = 1.40)$	Control $(n = 1.40)$	Control $(n = 1.40)$
	120)	120)	120)	140)	140)	140)
	$2  { m weeks}  (n$	4  weeks  (n	8 weeks $(n$	$2  { m weeks}  (n$	4  weeks  (n	8 weeks $(n$
	= 118)	= 120)	= 119)	= 137)	= 137)	= 140)
Never	0	0	0	47(34.3)	39(28.5)	34(24.1)
Occasionally	1(0.8)	0	0	25(18.2)	25(18.2)	25(17.7)
Once/week	1(0.8)	0	2(1.7)	7(5.1)	9(6.6)	10(13.5)
2-3/week	3(2.5)	3(2.5)	5 (4.2)	32(23.4)	40 (29.2)	44 (31.2)
4-6/week	8 (6.8)	6(5.0)	9 (7.6)	16(11.7)	17(12.4)	11 (7.8)
Daily	105 (89.0)	111 (91.7)	103 (86.6)	10(7.3)	7 (5.1)	8 (5.7)
Twice/day	75 (63.6)	83 (69.2)	87 (73.1)	2 (1.5)	0	0

Table 2 Parent-reported emollient application frequency at 2, 4 and 8 weeks (questionnaire data)

Data are n(%)

Table 3 Atopic dermatitis outcomes at 6 and 12 months

					Relative Risk
	Total	Intervention	Control	P-value	(95%  CI)
Primary					
outcome					
Cumulative	103/257	39/119	64/138	0.036	0.707 (0.516,
AD at 12	(40.1%)	(32.8%)	(46.4%)		0.967)
months					
Secondary					
outcomes					
AD at $12$					
months					
AD according to	76/253~(30%)	24/117~(20.5%)	52/136~(38.2%)	0.003	0.536(0.354,
the UK Working					0.813)
Party Diagnostic					
Criteria <sup>+</sup>					
AD at 6					
months		22 (122 (12 20)	10 /1 10 (05 00)	0.004	0 504 (0 005
AD according to	71/260 (27.3%)	22/120 (18.3%)	49/140 (35.0%)	0.004	0.524 (0.337, 0.012)
the UK Working					0.813)
Party Diagnostic					
Criteria '	79/960	99/190	F1 /140	0.009	0 502 (0 225
	(3/200 (39.107)	$\frac{22}{120}$	$\frac{31}{140}$	0.002	0.303 (0.323, 0.770)
AD	(20.1%)	(18.3%)	(30.4%)		0.779)

+ Point prevealance.

Abbreviations: AD = atopic dermatitis

## Figures

Figure 1 Trial profile

Figure 1 legend: \*Two infants in the control group were no longer eligible due to receiving phototherapy for jaundice after randomization. The primary outcome was the cumulative incidence of atopic dermatitis at 12 months.

Abbreviations: CW = Consent withdrawn, LTFU = Lost to follow-up

Figure 2 Point prevalence and cumulative incidence of AD at 12 months in the intervention and control groups

Figure 2 legend: The point prevalence of AD was calculated based on the number of infants in each group who met the UK Working Party Diagnostic Criteria for atopic dermatitis at 12 months (intervention: 24/117, control: 52/136), as assessed by the blinded investigator. The cumulative incidence of AD was calculated based on the number of infants diagnosed with AD at any point in the first 12 months (intervention: 24/117, control: 52/136).

Abbreviations: AD = atopic dermatitis

Figure 3 Kaplan-Meier plot of the proportion of in infants in the intervention and control group

without AD during the first 12 months of life

Figure 3 legend: The intervention group maintained AD-free skin for a longer period in the first 12 months than the control group (P = 0.016, log-rank test).

Abbreviations: AD = atopic dermatitis



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Figure 3\_Ní Chaoimh et al.