Mechanisms for *Alternaria alternata* Function in the Skin During Induction of Peanut Allergy in Neonatal Mice With Skin Barrier Mutations

Lauren M. Buelow, Akihiko Hoji, Kiet Tat, Lindsay M. Schroeder-Carter, Daniela J. Carroll and Joan M. Cook-Mills*

Departments of Pediatrics and Microbiology and Immunology, Herman B Wells Center for Pediatric Research, Indiana University School of Medicine, Indianapolis, IN, United States

Neonatal mice with heterozygous mutations in genes encoding the skin barrier proteins filaggrin and matrin (flaky tail mice [FT+/−]) exhibit oral peanut-induced anaphylaxis after skin sensitization. As we have previously reported, sensitization in this model is achieved via skin co-exposure to the environmental allergen *Alternaria alternata* (Alt), peanut extract (PNE), and detergent. However, the function of Alt in initiation of peanut allergy in this model is little understood. The purpose of this study was to investigate candidate cytokines induced by Alt in the skin and determine the role of these cytokines in the development of food allergy, namely oncostatin M (Osm), amphiregulin (Areg), and IL-33. RT-qPCR analyses demonstrated that skin of FT+/− neonates expressed Il33 and Osm following Alt or Alt/PNE but not PNE exposure. By contrast, expression of Areg was induced by either Alt, PNE, or Alt/PNE sensitization in FT+/− neonates. In scRNAseq analyses, Osm, Areg, and Il33 were expressed by several cell types, including a keratinocyte cluster that was expanded in the skin of Alt/PNE-exposed FT+/− pups as compared to Alt/PNE-exposed WT pups. Areg and OSM were required for oral PNE-induced anaphylaxis since anaphylaxis was inhibited by administration of neutralizing anti-Areg or anti-OSM antibodies prior to each skin sensitization with Alt/PNE. It was then determined if intradermal injection of recombinant IL33 (rIL33), rAreg, or rOSM in the skin could substitute for Alt during skin sensitization to PNE. PNE skin sensitization with intradermal rIL33 was sufficient for oral PNE-induced anaphylaxis, whereas skin sensitization with intradermal rAreg or rOSM during skin exposure to PNE was not sufficient for anaphylaxis to oral PNE challenge. Based on these studies a pathway for IL33, Areg and OSM in Alt/PNE sensitized FT+/− skin was defined for IgE induction and anaphylaxis. Alt stimulated two pathways, an IL33 pathway and a pathway involving OSM and Areg. These two pathways acted in concert with PNE to induce food allergy in pups with skin barrier mutations.

Keywords: anaphylaxis, keratinocyte, amphiregulin, oncostatin M, IL33, *Alternaria alternata*, peanut, food allergy
INTRODUCTION

Food allergy is a severe and potentially life-threatening disease that can develop early in life and affects about 8% of children (1–3). While there is no cure for food allergy, oral peanut immunotherapy is a promising therapeutic approach (4). The Learning Early About Allergy to Peanut (LEAP) trial identified that early introduction of peanut is beneficial in reducing development of peanut allergy in children (5, 6). However, although this intervention was efficacious, 9% of children were excluded from the study (7) because they were skin prick positive for peanut sensitization at the start of the study and another 5% were excluded because of reactions during oral peanut exposures (7). Notably, a number of these high-risk children who had early onset of peanut allergy also had eczema, suggesting a potential link between skin exposure to peanut and development of peanut allergy in children.

Food allergy occurs in up to 35% of children with atopic dermatitis (8). In fact, early-onset of atopic dermatitis (by age 2) is associated with increased risk of food allergy (9, 10). The development of atopic dermatitis is linked to loss-of-function mutations in the skin barrier genes SPINK5, a serine protease inhibitor, FLG (filaggrin), which crosslinks keratin fibers in keratinocytes (11) and Tmem79 (mattrin), a transmembrane protein involved in stratum corneum barrier function (12). These loss-of-function mutations in skin barrier genes have been found in infants and children with peanut allergy (13–17). Interestingly, in children, barrier defects caused by FLG mutations precede clinical eczema diagnosis (18). Similarly, in neonatal mice with heterozygous skin barrier mutations Flg and Tmem79 (Flaky Tail mice, FT+/-), skin sensitization to food allergens (19) occurs well before the spontaneous development of eczema (20), suggesting that allergen sensitization can occur before the onset of atopic dermatitis. Approaches are needed for clinical food allergy studies for stratifying individuals before their development of atopic dermatitis (18).

The development of atopic dermatitis and allergies is associated with exposure to environmental fungal allergens and house dust mite (HDM) allergen (21), commonly found in household dust (22–26). Studies have shown that patients with atopic dermatitis exhibit a higher frequency of IgE reactivity to HDM (27) or the fungal allergen Alternaria alternata (Alt) (28). Alternaria alternata is the most common fungi in mold contributing to asthma and the most common fungi known to be associated with allergic disease (29, 30). In a national survey of lead allergens in US homes, Alternaria antigens were found to be in 95–99% of dust samples at a concentration of 4.88 μg/g. Alternaria is very prevalent in homes in the US and can get into the household through hair, skin, clothing, pet fur, shoes, or through cracks in the house. The levels of Alternaria in the household can vary due to region, homeowner behavior, and housing characteristics but nonetheless Alternaria antigens were found in virtually all homes across the US (29).

Previously published data from our lab demonstrated that heterozygous skin barrier mutations in Flg and Tmem79 together with skin co-exposure to detergent, food allergen (egg or peanut), and a ubiquitous environmental allergen (HDM or Alt) at doses commonly found in household dust mediate the development of food allergy in neonatal mice (19). Since either HDM or Alt exposure with peanut induced food allergy, the induction of food allergy was not specific to fungal allergens. Although fungal allergens can exhibit protease activity, we have reported that Alt does not degrade the proteins in peanut extract (19). Moreover, the effect of Alt is not specific to peanut because co-exposure of Alt and chicken egg ovalbumin on skin of FT+/- pups also induce food allergy to OVA (19). The detergent on the skin functions to aid skin adsorption of saline and allergens (19). In our previous manuscript, Alt was required for the sensitization to food allergen, Alt/PNE co-exposure induced IL33 expression in the skin, and blocking the receptor for IL33 blocked anaphylaxis in neonatal mice. However, it is not known whether IL33 induction required co-exposure to Alt/PNE or whether Alt alone was sufficient in the FT+/- neonates. IL-33 is induced in another food allergy model that utilizes mechanical skin disruption with tape stripping that removes layers of epidermis (31). Also, anti-IL-33 antibodies block atopic dermatitis, at least in a model with 2, 4-dinitrochlorobenzene sensitization (32). Given the critical interplay between genetic alterations in skin barrier mutations and environmental allergen exposure for the development of food allergy, it is important to characterize the mechanisms for these interactions.

In skin with barrier defects, it is reported that amphiregulin (Areg) has a role in induction of itch (33) and in development of skin psoriasis (34). In atopic dermatitis in humans and mice, Areg is upregulated in several cell types in skin (35, 36). Also it is reported that Areg is induced in dendritic cells exposed to the environmental allergen Der p 1 in vitro (37). We have demonstrated that Areg is increased during allergic responses in the airways of neonatal mice (38). However, it is not known whether Areg plays a role during skin sensitization for induction of food allergy.

It is reported that Areg can be induced by oncostatin M (Osm) in tumor cells (39) and that the OSM receptor functions in induction of itch in atopic dermatitis (40, 41). The OSM receptor forms a complex with the IL-31 receptor A and then IL31 binding to this receptor drives itch in atopic dermatitis (40, 41). OSM in macrophages is upregulated by histamine, a product of mast cells (42). In the lung, OSM plays a role in airway remodeling, eosinophilic airway inflammation, and airway hyperresponsiveness (43), potentially by promoting mucosal epithelial barrier dysfunction (44). Transient transgenic overexpression of Osm in lung induces B cell activation, T cell differentiation, eosinophil accumulation, and aids in DC maintenance (45). However, it is not known whether OSM plays a role during skin sensitization for induction of food allergy. Thus, here we sought to investigate the relevance of IL33, OSM, and Areg in a model of food-induced anaphylaxis that relies on Alt sensitization via the skin. We demonstrated that in neonatal FT+/- mice, the environmental allergen Alt induces skin expression of IL33, Areg, and Osm which were required for generation of PNE-specific IgE and oral PNE-induced anaphylaxis.
MATERIALS AND METHODS

Mice

C57BL/6J mice, Fcer1a tm1Knt mice (FceR1a−/− mice) and Flaky Tail mice (FT−/− mice) with homozygous mutations in Flk b1f1/Tmem79 ma/m were from Jackson Laboratories, Bar Harbor, Maine. Mice were housed under specific pathogen-free barrier conditions. The Flaky tail mice on the C57BL/6 background have been previously described (12, 20, 46–48). All mice were randomly selected for the studies. The studies are approved by the Indiana University School of Medicine Institutional Review Committee for animals.

Allergens

Alternaria alternata extract (Alt, catalog #XPM1D3A2.5) was from Greer Labs. This is a standard extract from GREER used for patient immunotherapy. The protein profile in the Alt extract was assessed by SDS PAGE (Supplementary Figure 1). To generate peanut extract (PNE), peanuts (Trader Joe’s roasted and unsalted Peanuts) were ground and then 25 g were homogenized in 250 ml of 20 mM Tris buffered (pH 7.2) saline (TBS) (49). This was stirred for 2 h at room temperature and centrifuged at 3,000 g for 30 min. The aqueous middle layer was collected and centrifuged at 1,600 g for 45 min to remove residual particles and fat. The aqueous layer was collected and protein concentrations were determined by Pierce BCA Protein Assay Kit (Thermo Fisher Sci). Aliquots were stored at −20°C. The Alt and PNE extracts did not contain overt protease activity because co-administration of ALT and PNE did not alter the size of the major protein bands on SDS page chromatography (19).

Allergen Sensitization

(A) Treatment of offspring (19). FT−/− offspring of the mating of wild type (WT) C57BL/6 females with FT−/− males were confirmed by genotyping (19). FT−/−/Fcer1−/− offspring of the mating of FceR1a−/− (C57BL/6 background) females with FT−/−/Fcer1−/− males was confirmed by genotyping. At 3 days old, pups were gently taped four or five times on the back with 3M surgical hypoallergenic paper tape that does not disrupt the skin but gently removed dried skin after birth or hair after shaving (19).

(B) Intradermal administration of recombinant Areg and OSM. For studies with recombinant proteins, the pups received 5 skin sensitizations. For the pups treated with rAreg (catalog #554104, Biolegend) or rOSM (catalog #762806, Biolegend), the pups received an intradermal injection of 2.5 µg rAreg or rOSM just before each skin application of PNE. The doses for injection of these recombinant proteins were similar to those previous described for injection of mice (52–54).

(C) Neutralizing Antibody Blockade of Areg and OSM. For studies with neutralizing antibodies, the pups received 5 skin sensitizations. For the pups treated with goat IgG anti-Areg (catalog #AF989, R&D Systems), goat IgG anti-OSM (catalog #AF-495-NA, R&D Systems), or IgG isotype control (catalog #BD0083, Biocell) antibodies, the pups received an intraperitoneal injection of 10 µg antibody in saline at 1 h before skin sensitization #1, 12 µg antibody at 1 h before skin sensitization #2, 15 µg antibody at 1 h before skin sensitization #3, and then 29 µg antibody at 1 h before skin sensitization #4 and #5 to adjust for increases in body weight during neonate growth. Antibody doses were similar to those previous described for intraperitoneal injection of mice (33, 54, 55).

Oral Antigen Challenge, Anaphylaxis, and Tissue Analysis

At 48 h after the last allergen skin sensitization, the pups were weighed and baseline rectal temperatures of the pups were taken using a BAT-12 Microprobe Thermometer with a RET-4 thermocouple sensor type T rectal probe for neonatal mice (Physitemp Instruments Inc). Then, the pups received PNE by gavage (100 µl of 10 mg protein/ml sterile water) using a 24 gauge gavage needle (Pet Surgical, Agoura Hills, CA). Rectal temperatures were taken every 15–20 min for up to 80 min. The pups were placed back with the mothers. Then 4 h later, the pups were euthanized and tissues collected.

RNA Isolation and Quantitative Real-Time PCR

For skin collection from euthanized mice, the skin was sprayed with RNase ERASE between the shoulders at the sites of allergen exposure, and then circular skin punches were taken between the shoulders using an 8 mm skin puncher (Catalog # 33-37, Integra Miltex Disposable Biopsy Punch). The skin punches were placed in 2 ml collection tubes containing 100 µl of TriZOL (Life Technologies catalog # 15996018) and 50 µl of RNase ERASE before snap freezing in liquid nitrogen and stored in −80°C. Tissues were homogenized using Next Advance Bullet Blender Storm 24 Place Bead Homogenizer. Before homogenization, 500 µl TriZOL was added to Rino Bullet Blender Navy Bead Lysis Kit (Catalog # NAVYR5-RNA) and homogenized on level 12 for 4 min. 500 µl TriZOL was added again and beads were centrifuged at 4°C for 8 min at 1,200 rpm. Then, 200 µl chloroform (Catalog # BP1145, Fisher Scientific) was added to the lysates and tubes were placed on ice for 10 min. Skin tissue RNA was isolated using the Aurum Total RNA Fatty and Fibrous Tissue Kit (Catalog # 732-6830, Bio-Rad). The lysates were inverted periodically during 10 min on ice and then were centrifuged at 12,000 × g for 15 min at 4°C. The top aqueous layer was collected and added to an equal volume of 70%
Ethanol (made using RNA-free water (Catalog # BP561, Fisher Scientific) for 10 min on ice. The rest of the RNA extraction procedure followed the spin protocol of the Aurum Total RNA Fatty and Fibrous Tissue Kit for small tissue for mass > 10 mg (Catalog # 732-6830, Bio-Rad) on ice. A DNA digest was done by incubating the samples with 10 units of Benzonase Nuclease Ultrapure (Catalog # E8263-5KU, Sigma Aldrich) after the second wash of the DNA digest. RNA concentration, A260/280 ratio, and A260/230 ratio of each RNA sample was measured using a Take3 plate and Synergy H1 Microplate Reader. cDNA was generated using the Applied Biosystems High-Capacity cDNA Archive Kit. The RT-qPCR analysis was performed with a 7500 Fast Real Time PCR System (Applied Biosystems). Multiplex reactions were ran using TaqMan multiplex real-time PCR. Reactions were set up using Applied Biosystems QSY Probes (Catalog #4482777) and TaqMan Multiplex Master Mix (Catalog #4461882). Custom oligos (Applied Biosystems) were used from sequences obtained from NCBI Blast.

For scRNAseq, Cell Ranger UMI matrix outputs (provided by the genomic core) were analyzed by Seurat (v4.0.3) and https://dataview.ncbi.nlm.nih.gov/object/PRJNA728696?reviewer=tc2t0dsh81dc1okiftcmq6pri7 and https://dataview.ncbi.nlm.nih.gov/object/PRJNA728697?reviewer=4arak8qdugui8cujekvuqhpnp. The scRNAseq Analyses and Visualization

For scRNAseq analysis, Cell Ranger UMI matrix outputs (provided by the genomic core) were analyzed by Seurat (v4.0.3) (56). A standard pre-processing pipeline was performed prior to integration of WT and FT(+/−) transcriptomic data and subsequent generation of the unified UMAP was as described in Seurat vignettes. Curation of cluster-specific cell types on UMAPs were done first by running FindConservedMarkers to create a list of ranked and significant cluster-specific genes. Then, those cluster-specific genes were searched through web cell annotation resources including PanglaoDB (57) and Immgen (58) to identify the most fitted cell types of each clusters. This curation process was further refined by comparing and contrasting our gene list to those skin scRNAseq data from similar publications (59, 60). In addition to Seurat visualization, ggplot2 (v.3.3.2) (61) was used to generate UMAPs with overlays of OSM, Areg, and IL33.

PNE-Specific Antibody, MCPT-1, OSM, Areg, and IL33 ELISAs

Serum and plasma was collected at 4 h after oral gavage with PNE. Serum anti-peanut IgE, IgG1, and IgG2a were determined by ELISA. High binding 96 well EIA/RIA plate (Fisher Scientific, catalog #50-823-480) were coated with 50 µl of 10 µg PNE protein/ml in carbonate buffer overnight at 4°C. Plates were washed with PBS-0.05%TWEEN and blocked with PBS + 3% BSA for 2h at room temperature. Plates were washed with PBS-Tween. Serum was plated in PBS + 1% BSA at a 1/10 dilution for IgE, 1/1,000 for IgG2b and 1/40,000 for IgG1 analysis and then incubated overnight at 4°C. Plates were washed with PBS-0.05%TWEEN and incubated for 1 h at room temperature with 100 µl of 2 µg/ml biotin-labeled secondary antibody in PBS-1% essentially immunoglobulin-free BSA. Plates were washed with PBS-Tween and incubated with 100 µl/well streptavidin-HRP (1:1,000 dilution in PBS, GE Healthcare, catalog #RPN1231-2ML). Plates were washed in TBS-Tween and incubated with 100 µl TB substrate (eBioscience catalog#00-4201-56, 3,3’,5,5’-Tetramethylbenzidine; HRP substrate). At the indicated times in the figures, the reaction was stopped with 100 µl 2M phosphoric acid and plates read at 450 nm in a microtiter plate reader. MCPT-1 in plasma was measured by ELISA according to manufacturer’s instructions (catalog #50-174-16, Fisher Scientific). Serum IL33, OSM and Areg were analyzed by ELISA according to manufacturer’s instructions (IL33 Cat #DY3626-05 R&D Systems, OSM Cat #DY495-05 R&D Systems, Areg Cat#DY989 R&D Systems, MCPT1 Cat#88-7503 Invitrogen).

Genotyping for Flg<sup>ft</sup> and Tmem79<sup>ma</sup>

Offspring of the mating of wild type C57BL/6 females with Flg<sup>−/−</sup>/Tmem79<sup>ma/ma</sup> males were confirmed by genotyping. Briefly, for Flg PCR detection, murine genomic DNA was isolated from tail clips using the Quantabio Extractna DNA Prep kit (catalog #950901) and amplified with the primers: Forward: 5’-CATCTCCAGTCAGGCGTACC-3’, Reverse: 5’-GCTGCCGTGCGCCAGACTCG-3’.

PCR amplification conditions were as follows: (96°C, 5 min) x1; (96°C 30 s, 60°C 30 s, 72°C 30 s) x35; (72°C, 5 min) x1. Amplified transcripts were...
then digested with the restriction enzyme Acc I (New England Biolabs, catalog #R01611s) yielding either a wild type FLG band at 678 bp or two fragments of the mutant FLG at 559 and 134 bp as determined by gel electrophoresis.

**Statistics**

Data were analyzed by a one way ANOVA followed by Wilcoxon’s non-parametric multiple comparisons test (SigmaStat, Jandel Scientific, San Ramon, CA). Presented are the means ± the standard errors. The statistical analysis for the change in temperature was done using the area of the curve by summation of the temperature changes at 15, 30, 45, 60, and 80 min for each pup. The data in the figures include both sexes of the offspring because there were no differences in food allergy outcomes by sex as previously described (19).

**RESULTS**

**Alt Increases Expression of IL33, Areg, and Osm in Skin of FT+/− Neonatal Mice**

We previously demonstrated that co-exposure to detergent, Alt, and PNE induce peanut allergy in FT+/− neonatal mice, but not in WT neonatal mice (19). Because the function of Alt in induction of food allergy is not known, we addressed whether Alt induced expression of OSM, Areg, and IL33, which have roles in atopic dermatitis (19, 32, 33, 35, 36) and allergies in the lung (44). We performed quantitative real-time PCR (RT-qPCR) analysis for Osm, Areg, and IL33 in the skin of neonatal mice at 4 h after the last skin sensitization on postnatal day 16 (Figure 1A). Osm and IL33 gene expression was significantly increased in the Alt and Alt/PNE sensitized FT+/− neonates as compared to the saline-treated FT+/− neonates (Figures 1B,D), but not in the PNE only sensitized skin. In contrast, gene expression of Areg was induced in all three treatment groups (Alt, PNE, and Alt/PNE) (Figure 1C) as compared to saline-treated FT+/− neonates. There was no increase in expression of these genes following skin sensitization of WT neonates (Figures 1B–D).

To determine which cells expressed IL33, Osm, and Areg in the Alt/PNE-exposed skin of FT+/− neonates and WT neonates, single cell RNAseq and UMAP analyses was performed (Figure 2) using skin biopsies from neonates at 4 h after the last skin sensitization (Figure 1A). Although eosinophils are present in the skin in this model as determined by histology (19), eosinophils were not detected in single cell RNAseq analyses of the skin, as has been a difficulty in the field for single cell RNAseq analyses of tissue eosinophils. The cell types in the UMAP clusters (Figure 2A) were assigned by gene expression profiles (Supplementary Table 1). Of note, cluster 6 keratinocytes had a larger number of cells in the Alt/PNE exposed FT+/− as compared to Alt/PNE exposed WT neonates (Figure 2A, red circle). The cluster 6 keratinocytes of the FT+/− neonates also expressed Osm, Areg, and IL33 (Figures 2B,C), indicating a source of these proteins not present in the WT neonates. The cell types expressing OSM that were common to Alt/PNE-treated FT+/− and WT skin were macrophages (clusters 1,2), neutrophils (clusters 4,12), mast cells (cluster 20), basophils (cluster 22), and dendritic cells (cluster 5) (Figures 2B,C). The cell types expressing Areg that were common to FT+/− and WT with Alt/PNE were T cells (cluster 9) and macrophages (cluster 1) (Figures 2B,C). The cell types expressing IL33 that were common to FT+/− and WT with Alt/PNE were fibroblasts (cluster 14) (Figures 2B,C). Thus, several cell types expressed Osm, Areg, and IL33 but a distinction for the Alt/PNE-treated FT+/− neonates was expression of these cytokines by keratinocytes in cluster 6.

**Areg and OSM Are Required but Not Sufficient for the Induction of Peanut Allergy in FT+/− Neonatal Mice**

In Figure 1, we showed that IL33, Areg, and Osm gene expression was induced following Alt/PNE skin sensitization of FT+/− neonates. Previously we reported that blocking the IL-33 receptor ST2 with anti-ST2 antibodies blocks generation of PNE-specific IgE and PNE-induced anaphylaxis (19). To address a role for IgE in this food allergy model, oral PNE-induced anaphylaxis was blocked in Alt/PNE skin sensitized FceR1a−/− FT+/−/− pups (Figures 3A,B) that had serum PNE-specific IgE (Figure 3C). Then, we determined whether Areg and OSM are required for the induction of peanut allergy. To test this, neutralizing anti-Areg or anti-OSM antibodies were administered at previously described doses (33, 54, 55) by intraperitoneal injection at 1 hr before each skin sensitization with Alt/PNE as in Figure 3D. Oral PNE challenge induced anaphylaxis in Alt/PNE sensitized pups (Figures 3B,E). As expected (19), neonates had a reduced ability to control temperature over the time course as compared to adults (62–65), but temperatures begin to rebound after 105 min and then pups are placed back with their mothers (19). Administration of neutralizing anti-Areg or anti-OSM during Alt/PNE sensitization of FT+/− neonates blocked oral PNE-induced anaphylaxis (Figure 3E). Together, our previous observations with anti-ST2 blockade (19) and our current findings with anti-Areg or anti-OSM suggest that ST2, OSM, and Areg are required for the induction of food allergy.

It was also determined whether intradermal recombinant (r) OSM, rAreg, or rIL-33 during skin co-exposure with PNE was sufficient to bypass the need for Alt for induction of food allergy as measured by oral PNE-induced anaphylaxis. rOSM, rAreg, or rIL-33 was administered at a concentration of 2.5 μg/neonate by intradermal injection (52–54) and immediately followed by topical application of detergent and PNE as in the timeline in Figure 3D. Skin sensitization with rIL-33/PNE was sufficient to induce anaphylaxis following oral challenge with PNE (Figure 3E). In contrast, rOSM/PNE and rAreg/PNE were not sufficient to induce anaphylaxis following oral challenge with PNE (Figure 3E), suggesting that OSM and Areg are necessary but not sufficient to induce anaphylaxis following oral challenge with PNE.

**IL33, OSM, and Areg Regulate Induction of Anti-PNE IgE and Mast Cell Degranulation**

Next we addressed whether these cytokines regulate IgE and mast cell degranulation in this model because anaphylaxis...
FIGURE 1 | Allergen skin sensitization of FT+−/− neonates induced skin expression of IL33, Areg, and Osm. (A) Timeline for mating and for FT+−/− pup treatments with saline, Alt, PNE, and Alt/PNE allergens. Tissues were collected 4 h after the last pup skin sensitization. Relative expression of skin expression of (B) Osm, (C) Areg, and (D) IL33 as determined by RT-qPCR. N = 8–10/group. The data in all figures include both genders for pups because there were no differences in outcomes by gender as we previously reported for Alt/PNE skin sensitization (19). *p < 0.05 as compared to the saline skin-sensitized groups or WT groups. Alt, Alternaria alternata extract; Areg, amphiregulin; FT−/−, flaky tail mice homozygous for filaggrin and mattrin mutations; FT+/−, flaky tail mice heterozygous for filaggrin and for mattrin mutations; IL33, interleukin-33; Osm, oncostatin M; PND, postnatal day; PNE, peanut extract; WT, wild type.

was dependent on FceR1 (Figure 3B). FceR1 is expressed by mast cells, and we have reported that mast cells degranulate in skin and intestine of Alt/PNE-sensitized FT+−/− pups (19). At 4 h post oral gavage as in the timeline in Figure 3D, Alt/PNE, but not Alt or PNE alone, induced anti-PNE-specific IgE, IgG2b, and IgG1 (Figures 4A–C). The increase in Alt/PNE-induced anti-PNE-specific IgE was inhibited by anti-OSM neutralizing antibody, whereas there was no effect of the anti-Areg neutralizing antibody or the isotype control antibody (Figures 4A,B). However, the Alt/PNE-induced anti-PNE-specific IgG1 was not altered by anti-OSM, anti-Areg, or isotype antibodies (Figure 4C). Interestingly, rOSM/PNE, rAreg/PNE, or rIL33/PNE was sufficient to induce anti-PNE-specific IgE and IgG2b (Figures 4A,B). Anti-PNE-specific IgG1 was induced by rOSM/PNE or rAreg/PNE but not by rIL33/PNE (Figure 4C). Thus, our current findings together with our previous observations that anti-ST2 neutralizing antibodies block generation of anti-PNE-specific IgE (19) suggest that OSM and the IL33 receptor ST2 were required for the production of anti-PNE-specific IgE.

As a measure of mast cell degranulation in these studies, the mast cell granule protein MCPT1 was determined in the serum at two time points, 4 h after skin sensitization and 4 h after oral PNE. At 4 h after oral PNE (Figure 4D), Alt/PNE-skin-sensitized and Alt-skin-sensitized FT+−/− pups had increased serum MCPT1 as compared to PNE-skin-sensitized FT+−/− pups. This increase in MCPT1 in the Alt-skin-sensitized FT+−/− pups occurred even though the Alt-skin sensitized pups did not have elevated anti-PNE specific antibodies (Figures 4A–C) and did not have oral PNE-induced anaphylaxis (Figure 3E). This suggests that Alt induces mast cell degranulation without IgE and that MCPT1 release was not sufficient for anaphylaxis in Alt-sensitized pups. Furthermore, at 4 h after the last skin sensitization (without oral gavage with PNE), there was also increased serum MCPT1 (Figure 4E). The absolute amounts of MCPT1 after skin sensitization are not directly compared with...
FIGURE 2 | UMAP visualization of skin scRNAseq analysis. (A) UMAPs of scRNAseq from skin punches of FT+/− and WT neonates treated with Alt or Alt/PNE with cluster-specific cell type annotation. (B) UMAP projection of skin cells expressing Osm (green), Areg (blue), and Il33 (red) above the threshold expression value (Seurat logNormalized) > 1.5. Annotated cell type designation to numbered clusters in (A) is listed as follows: Dendritic, dendritic cells; Endo, endothelial cells; Fib, fibroblasts; ILC, innate lymphoid cells; KC, keratinocytes; Mac, macrophages; PMN, neutrophils; SM, smooth muscle cells; Stem, stem cells. (C) Cell clusters with abundant gene expression of Osm, Areg, and Il33.
FIGURE 3 | Areg and OSM during co-exposure to PNE was necessary but not sufficient for FceR1-dependent oral PNE-induced anaphylaxis in FT+/− neonatal mice. (A) FT+/− FceR1a−/− males were mated with FceR1a−/− females. The pups (FT+/− FceR1a−/−) were skin sensitized four times as in the Timeline. (B) Oral (Continued)
Pathways for Alt/PNE Induction of Skin IL33, OSM, and Areg Gene Expression

Oral PNE-stimulated anaphylaxis was induced in pups skin-sensitized with Alt/PNE but not PNE alone, Alt alone (Figure 3E) or saline (19). To determine pathways by which Alt induces expression of Il33, Osm, and Areg, RT-qPCR was performed to analyze expression of these genes in the skin of neonatal mice treated as in Figure 3D. In Figure 5A, expression of Osm was induced by Alt and Alt/PNE. The induction of Osm expression by Alt/PNE was not blocked by anti-OSM or anti-Areg and not induced by rOSM/PNE, rAreg/PNE, or rIL33/PNE. In Figure 5B, expression of Areg was induced by Alt, PNE, Alt/PNE, or rOSM/PNE, but not rIL33/PNE or rAreg/PNE. Taken together, the data suggest that OSM induces Areg in these Ft+/− neonates. The induction of Areg expression by Alt/PNE was blocked by anti-OSM but not by the isotype control (Figure 5B), further supporting that OSM induces Areg. The inhibition of Areg gene expression by anti-Areg co-administration with Alt/PNE suggests that Areg is regulated in an autocrine manner, consistent with other descriptions of Areg autocrine loops (39).

In Figure 5C, Il33 expression was induced by Alt and Alt/PNE. This Alt/PNE-induced Il33 expression was not blocked by anti-OSM or anti-Areg (Figure 5C). Also Il33 expression was not induced by rOSM/PNE or rAreg/PNE (Figure 5C), indicating that Alt induced a pathway with Il33 and another pathway with OSM and Areg. Furthermore, putting the data together with data from Figures 3E, 5, it suggests that the rIL33/PNE was sufficient to sensitize for oral PNE-induced anaphylaxis through a combined pathway of Il33 and PNE-stimulated Areg. Also, although rOSM/PNE and rAreg/PNE induced production of anti-PNE-specific IgE (Figure 4A), these skin sensitizations were not sufficient for oral PNE-induced anaphylaxis (Figure 3E), suggesting that IL33-mediated signals were required to induce a state for responsiveness to oral PNE-induced anaphylaxis.

OSM, Areg, and IL33 Are Elevated in Plasma During Alt/PNE Induced Food Allergy

To investigate whether there were systemic plasma levels of OSM, Areg, and IL-33 in the pups from Figures 3D,E, plasma levels of OSM, Areg, and IL-33 in Ft+/- pups were determined by ELISAs. Group comparisons were made to pups exposed to PNE or pups exposed to Alt/PNE/anti-Areg since they did not have anaphylaxis. Plasma OSM was elevated for neonates that were skin sensitized with Alt or Alt/PNE as compared to PNE sensitization (Figure 6A). This increase in plasma OSM for Alt/PNE-skin-sensitized neonates was not altered by anti-OSM, anti-Areg, or isotype antibodies (Figure 6A). Plasma OSM was not increased in neonates with rAreg/PNE or rIL33/PNE (Figure 6A). Interestingly, intradermal rOSM injections during skin sensitization with PNE increased plasma OSM even at 2 days after the injections, albeit not to the levels induced by Alt/PNE skin sensitizations (Figure 6A). Plasma Areg was reduced in neonates administered anti-OSM or anti-Areg but not isotype antibody during Alt/PNE-skin-sensitization (Figure 6B). The inhibition by anti-Areg is consistent with autocrine amplifying functions reported for Areg (39). Skin sensitization with rOSM/PNE, rAreg/PNE and rIL33/PNE increased plasma Areg protein concentrations, similar to concentrations induced by PNE alone (Figure 6B). This is consistent with PNE induction of skin Areg expression shown in Figure 5B. Plasma IL33 was elevated in neonate skin sensitized with Alt or Alt/PNE compared to PNE alone (Figure 6C). Plasma IL33 was slightly increased in rOSM/PNE-sensitized pups, but anti-OSM antibodies did not significantly reduce Alt/PNE-induced IL33 expression (Figure 6C). The increase in plasma IL33 in neonates sensitized with Alt/PNE was blocked by anti-Areg antibodies. However, rAreg/PNE did not increase plasma IL33.

DISCUSSION

We demonstrated that oral PNE-induced anaphylaxis in Alt/PNE-skin-sensitized Ft+/- pups was dependent on FceR1. Alt stimulated an increase in expression of Il33, Osm, and Areg in the skin of Ft+/- but not WT neonates. This Alt-induced expression of Il33, Osm, and Areg functioned in concert with food allergen on the skin of Ft+/- neonatal mice to sensitize neonates to food allergen, such that oral food allergen-challenge induced anaphylaxis. A model for Alt-stimulated pathways during the induction of food allergy is depicted in Figure 7. Briefly for Ft+/- neonates, topical PNE
FIGURE 4 | IL33, OSM, and Areg regulate induction of serum anti-PNE IgE, anti-PNE IgG2b, anti-PNE IgG1, and MCPT1. Serum was from pups treated as in Figure 3E. At 4 h after oral PNE-gavage, serum anti-peanut specific (A) IgE, (B) IgG2b, and (C) IgG1 were measured by ELISA. (D) Serum MCPT-1 at 4 h after oral gavage with PNE was measured by ELISA. (E) Serum MCPT-1 at 4 h after the last skin sensitization from pups treated as in Figure 3D was measured by ELISA. *p < 0.05 as compared to the PNE only skin-treated group. Alt, Alternaria alternata extract; Areg, amphiregulin; IL33, interleukin 33; OSM, oncostatin M; PNE, peanut extract.
FIGURE 5 | Skin Il33, Osm, and Areg gene expression in skin-sensitized FT+/− pups. Pups were treated as in Figure 3D, but with the addition of a group receiving saline on the skin. Skin was collected at 4 h after the oral PNE gavage and analyzed by RT-qPCR for expression of (A) Osm, (B) Areg, and (C) Il33. N = 6–10/group. *p < 0.05 as compared to the saline skin-treated group. Also, for skin OSM mRNA (A), the Alt/PNE/anti-Areg Ab group is not significantly different than the Alt/PNE group (p = 0.69) or the Alt/PNE/isotype Ab group (p = 0.39). Alt, Alternaria alternata extract; Areg, amphiregulin; IL33, interleukin 33; OSM, oncostatin M; PNE, peanut extract.
FIGURE 6 | OSM, Areg and IL33 are elevated in plasma during Alt/PNE induced food allergy. Plasma was collected at 4 h after oral PNE gavage (treated as in Figure 3D). Plasma (A) OSM, (B) Areg, and (C) IL33 were measured by ELISA. N = 8–10/group. (A,C) *p < 0.05 as compared to the PNE only skin-treated group. (B) *p < 0.05 as compared to the anti-OSM/Alt/PNE treated group. Alt, Alternaria alternata extract; Areg, amphiregulin; IL33, interleukin 33; OSM, oncostatin M; PNE, peanut extract.
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FIGURE 7 | Model for skin cytokine network in Alt/PNE induction of food allergy. Illustrated is a working model for a network of expression of IL33, OSM and Areg during Alt/PNE treated FT+/− skin. Alt induced two pathways, an IL33 pathway and an OSM/Areg pathway in the FT+/− skin. These pathways mediate induction of anti-PNE IgE when FT+/− skin is sensitized by co-exposure of Alt with PNE antigens. The data in Figure 5B also indicated an autocrine function for Areg in the FT+/− skin. Furthermore, PNE alone on the skin induced expression of Areg. When FT+/− skin is sensitized with rIL33/PNE, the IL33 pathway, the PNE-induced Areg pathway and the PNE antigens are sufficient for sensitization and oral-PNE-induced anaphylaxis.

Skin of FT+/− neonates:

- Alt
- PNE
- OSM
- IL33
- Areg
- keratinocyte
- macrophage
- neutrophil
- basophil
- mast cell
- dendritic cell
- keratinocyte fibroblast
- keratinocyte
- macrophage
- T cell
- Areg can induce Keratinocyte proliferation

Anti-PNE IgE FcεR1

ST2

Oral PNE-induced anaphylaxis in FT+/− neonates

alone on the skin induced expression of Areg but not IL33 or Osm. However, Alt induced skin expression of Osm, Areg, and IL33. The single cell RNAseq analyses of the skin indicated that Osm, Areg, and IL33 were expressed by several cell types. Of note, there were more cells in keratinocyte cluster 6 that expressed OSM, IL33, and Areg in FT+/− neonates than WT neonates at the site of Alt/PNE-epicutaneous exposure. Further ongoing studies will address cluster 6 keratinocyte generation and function. In antibody blocking studies, anti-OSM inhibited Alt/PNE-induced skin expression of Areg but anti-Areg did not inhibit Alt/PNE induced Osm expression. Anti-Areg and anti-OSM antibodies did not inhibit Alt/PNE-induced expression of IL33, indicating that Alt induces both an IL33-dependent pathway and an OSM/Areg-dependent pathway. Anti-Areg antibodies
reduced Areg expression suggesting an autocrine Areg pathway, consistent with reports in other inflammatory processes that OSM can induce an Areg autocrine function. For example, OSM induces an Areg autocrine loop during osteoclast differentiation (39). The OSM/Areg pathway and the IL33/ST2 pathways are both important for oral PNE-induced anaphylaxis in Alt/PNE-sensitized neonates because anaphylaxis is blocked by anti-ST2 (the IL33 receptor) (19), anti-Areg, or anti-OSM antibodies. rIL33/PNE was sufficient for sensitization and oral PNE-induced anaphylaxis, since rIL33 bypassed the need for Alt induction of IL33 and the skin exposure to PNE induced Areg expression. This suggested that the co-activation of an IL33 pathway and Areg pathway was sufficient to induce food allergy in the presence of food allergens. The lack of anaphylaxis in the rOSM/PNE or rAreg/PNE sensitized neonates suggests that an OSM/Areg pathway was not sufficient. However, an OSM/Areg pathway was required for anaphylaxis since administration of anti-OSM or anti-Areg antibodies during sensitization with Alt/PNE blocked oral PNE-induced anaphylaxis. With regards to anti-PNE-specific IgE, it was induced in FT+/- neonate skin exposed to rOSM/PNE, rAreg/PNE, or rIL33/PNE. The Alt/PNE-induced generation of anti-PNE specific IgE was blocked by anti-OSM but not anti-Areg antibodies and, in our previous report, generation of anti-PNE specific IgE was blocked by anti-ST2 (19). This suggests that OSM, IL33, and PNE are needed for generation of anti-PNE-specific IgE. OSM overexpression has been reported to have a role in activation of B cells, at least in lung inflammation, but antibody production was not examined (45). Since anti-OSM blocking antibodies inhibited generation of anti-PNE IgE but not anti-PNE IgG2b or IgG1, it suggests that OSM may have a unique role in IgE production after allergen skin sensitization of FT+/- pups.

Allergen-specific IgE does not always predict whether allergen will induce anaphylaxis. In humans, it is reported that food-specific IgE in sensitized patients does not always associate with anaphylaxis upon oral food challenge (67). Similarly, in the FT+/- pups, with different skin stimuli, anti-PNE IgE did not always associate with oral PNE-induced anaphylaxis. Briefly, oral PNE induced anaphylaxis in Alt/PNE-sensitized pups and this was dependent on FceR1 which binds IgE. Also, oral PNE induced anaphylaxis in the pups skin sensitized with rIL33/PNE. In contrast, anaphylaxis was not induced by oral PNE in pups skin sensitized with rAreg/PNE or rOSM/PNE, even though these pups had serum anti-PNE-specific IgE and increased serum MCPT1 after oral PNE gavage. Therefore, although IgE binds to the FceR1 on mast cells and basophils (68), the anti-PNE-specific IgE was not sufficient for induction of anaphylaxis in the absence of Alt stimulation of mediators such as IL33. IL33 can effect several cell types, including reports that it stimulates mast cell expansion (69).

In the rAreg/PNE or rOSM/PNE sensitized FT+/- pups which had increased anti-PNE specific IgG1, the IgG1 may have limited the oral PN-induced anaphylaxis because IgG1 binds to FcγRIIB which functions to limit the acute effector functions of mast cells and basophils (68). Furthermore, in the rIL33/PNE skin sensitized FT+/- neonates, there was not an increase in anti-PNE specific IgG1 and these pups had the greatest drop in temperature upon oral gavage. This suggests that in the absence of induction of anti-PNE-specific IgG1 in the rIL33/PNE sensitized neonates, there may be increased acute IgE-mediated signals for anaphylaxis.

Mast cell degranulation is induced by antigen crosslinking of IgE on FceR1, resulting in release of granule contents and increases in serum MCPT1 (70, 71). Besides FceR1/IgE, multiple other stimuli activate mast cells and can induce differential release granule and non-granule mediators from mast cells (72–75). Furthermore, basophil mediators may also be involved in anaphylaxis as the FcεR1 receptor is also expressed on basophils (68).

We have previously reported that oral PNE induces an increase in serum MCPT1 in Alt/PNE sensitized FT+/- pups and that this is blocked by anti-ST2 (19), suggesting that IL33 is necessary. Our previous publication also demonstrated by histology that in FT+/- neonates, mast cell degranulation occurs both in the skin at 4 hrs after Alt/PNE skin sensitization and 2 days later at 4 hrs after oral gavage with PNE in the intestine of the Alt/PNE-sensitized pups (19). This suggests that MCPT1 is released from mast cells both after skin sensitization and again after oral PNE challenge. The measurement of MCPT1 at 4 h is similar to the time frame for maximal serum MCPT1 in a study with intraperitoneal MCPT1 injections (76). Their study also showed that serum MCPT1 was cleared at about 24 h but that was from a single injection (76). A limitation of our current study is that it is not known whether systemic MCPT1 levels released after skin sensitization were sustained for 48 hrs and contributing to the systemic levels measured after oral PNE challenge. This is being addressed in ongoing studies of systemic mediators. Nevertheless, we now demonstrate that serum MCPT1 was also increased in FT+/- neonates sensitized with rOSM/PNE, rAreg/PNE, rIL33/PNE, or Alt as compared to PNE alone when examined 4 h after the last skin sensitization and at 4 hrs after oral PNE. Although both an IL33 pathway and an OSM/Areg pathway were sufficient for increased serum MCPT1 in this model, neutralizing antibodies for each of these cytokines did not block the increase in serum MCPT1. Together these data suggest that there is a redundancy in activation of MCPT1 release by the IL33 and the OSM/Areg pathways. It is has been reported that inhibition of Alt protease activity blocks Alt-induced increases in MCPT1 levels in the lung (77). In the food allergy model with FT+/- mice, we previously reported that PNE sensitization occurs with skin co-exposure to Alt/PNE or HDM/PNE, and the Alt extract did not contain protease activity, at least not toward PNE proteins (19, 66). Further studies to define how Alt activates OSM, Areg, IL33, and mast cell degranulation in FT+/- neonate skin are under investigation in our lab and beyond the scope of this report. Nevertheless, our data indicated that skin sensitizations with these recombinant cytokines and PNE were sufficient for mast cell MCPT1 release in FT+/- neonates but not WT neonates.
We also demonstrated that Alt skin sensitized FT\(^{+/−}\) pups had an increase in plasma levels of Areg, OSM, and IL33 protein, whereas FT\(^{+/−}\) pups that were skin sensitized with PNE did not have elevated plasma OSM or IL33. Thus, these mediators have systemic effects that may regulate activation of responses in distant sites. Intradermal administration of rAreg, rOSM, or rIL33 with topical PNE increased plasma Areg, OSM, and IL33, respectively. A systemic effect of mediators from the skin would be consistent with skin-derived factors blocking oral allergen induction of tolerance. For example, we reported that in FT\(^{+/−}\) pups, oral PNE pre-exposure before skin sensitization induces tolerance to oral PNE (19). However, this tolerance to PNE was blocked when skin was simultaneously exposed to Alt (19), suggesting that Alt-induced signals from the skin are systemic in that they limit induction of tolerance to oral peanut in the gut. This may have implications for children because there are clinical trials aimed to induce tolerance using oral peanut, but the skin of these children are also exposed to dust that contains Alt. Systemic effects of OSM, Areg, and IL33 are currently under investigation in our lab.

It has not been previously reported that Osm and Areg are present in the skin during induction of food allergy. However, Osm gene expression has been described in the lung in allergic diseases and in the skin with atopic dermatitis (54, 78). It is reported that in humans, there is an association of increased Osm gene expression in blood cells with anaphylaxis to any allergen (79). However, the function or mechanism for induction of Osm gene expression was not described. It has been reported that OSM mRNA and protein are increased in nasal passages and lung of patients with chronic rhinosinusitis and that OSM treatment of epithelial cells at air-liquid interface reduces barrier function (44), but the mechanisms for induction of Osm are not known. OSM induces IL33 expression in mouse liver endothelial cells (80) and overexpression of OSM induces mouse lung IL33 expression (81). However, in skin of FT\(^{+/−}\) neonates, rOSM/PNE did not induce IL33 gene expression and neutralizing anti-OSM antibodies did not reduce Alt/PNE-induced IL33 gene expression. This suggests that there may be differences in pathways for induction of IL33 expression by different tissues or stimuli (78). Reports indicate that keratinocytes produce IL33 during psoriasis (82) and in the Alt/PNE-stimulated skin of FT\(^{+/−}\) neonates, IL33 was primarily expressed by keratinocytes and fibroblasts.

Areg has been studied in psoriatic skin (34). It is also reported that skin keratin induces expression of Areg that induce thymic stromal lymphopoietin (TSLP)-mediated activation of itch (33). In osteoclasts in the bone, there is an Areg autocrine loop induced by OSM during osteoclast differentiation (39). Similarly, our data suggests that Alt stimulates an Areg autocrine loop during Alt/PNE stimulation of FT\(^{+/−}\) skin. The mechanism for Alt stimulation of keratinocytes in the skin of FT\(^{+/−}\) neonates may be through TLR2/4 since Alt binds and activates TLR2/4 (83) and these receptors are expressed by keratinocytes, fibroblasts and leukocytes (84, 85). Alt, which can have protease activity, does not alter the molecular weight of the major peanut proteins (Ara h 1, Ara h 3 acidic subunits, Ara h 3 basic subunit, Ara h 2, and Ara h 6) (19), albeit this does not preclude potential Alt protease effects on proteins in the skin.

Interestingly, in addition to a role for IL33 in initiation of food allergy (19), IL33 plays a role in re-challenge with a food allergen. This was demonstrated a study where anti-IL33 improved desensitization in peanut-allergic participants (86). As a further link between IL33 in food allergy and atopic dermatitis, patients with atopic dermatitis have a 10-fold increase in IL33 as compared to healthy controls (87). In our studies, the FT\(^{+/−}\) neonatal mice (without allergen stimulation) did not have elevated skin expression of IL33 at baseline as compared to WT pups. Moreover, the FT\(^{+/−}\) pups did not develop atopic dermatitis until a few months old (20). Moreover, in the FT\(^{+/−}\) pups, Alt but not PNE stimulated an increase in IL33 and this occurred well before FT\(^{+/−}\) pups spontaneously develop atopic dermatitis. Thus, because FT\(^{+/−}\) pups can develop atopic dermatitis at a few months old without food allergy and because food allergy can develop before atopic dermatitis, it suggests that with these skin barrier mutations in FT\(^{+/−}\) mice, food allergy or atopic dermatitis can develop independently.

In conclusion (Figure 7), PNE on the skin of FT\(^{+/−}\) neonates induced Areg, but did not induce Osm, IL33, anti-PNE-specific IgE, or oral PNE-induced anaphylaxis. In contrast, Alt induced two pathways, an IL33 pathway and an OSM/Areg pathway. Although IL33, Osm, and Areg expression occurred in several cell types in Alt/PNE-stimulated FT\(^{+/−}\) neonate skin, there was a keratinocyte cluster, expressing Osm, IL33, and Areg, that was not present in Alt/PNE-stimulated skin of WT neonates (Figure 2). Since the effect of Alt only occurred in skin of FT\(^{+/−}\) but not WT neonates, it suggests that keratinocyte function is altered in the FT\(^{+/−}\) neonates. Functionally, the Alt-induced Osm and IL33 participated in generation of anti-PNE-specific IgE and IgG2b in FT\(^{+/−}\) neonates. An IL33 pathway and an OSM/Areg pathway mediated activation of mast cells as demonstrated by elevated serum MCPT1 in FT\(^{+/−}\) neonates. Also OSM was required for the Alt-induction of Areg in FT\(^{+/−}\) neonates. The data also indicate that there is an Areg autocrine loop. Areg may function in induction of keratinocyte proliferation since Areg is reported to induce keratinocyte proliferation (55) (Figure 7). In addition to local effects in the skin, FT\(^{+/−}\) skin exposure to Alt generated systemic levels of Areg, OSM, and IL33 as these were increased in the plasma. Importantly, OSM, Areg, and IL33 functioned in Alt/PNE sensitization of FT\(^{+/−}\) neonates for generation of sensitivity to oral PNE-induced anaphylaxis. Interestingly, the Alt induction of IL33, Areg and OSM occurred in neonates well before the development of atopic dermatitis in the FT\(^{+/−}\) mice. These data have important implications for studies of detection of risks for sensitization to food allergens.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: NCBI SRA BioProject, accession numbers: PRJNA728696 and PRJNA728697 (private).
ETHICS STATEMENT

The animal study was reviewed and approved by Indiana University School of Medicine Institutional Review Committee for animals.

AUTHOR CONTRIBUTIONS

LB participated in performing and interpreting experiments, data and statistical analysis, and manuscript preparation. AH analyzed the RNaseq data and generated figures, as well as participated in performing and interpreting experiments and participated in manuscript preparation. KT did the experiments for RNaseq analyses. LS-C participated in performing and interpreting experiments. DC participated in data interpretation and manuscript preparation. JC-M conceived of the study design and participated in performing experiments, statistical analysis, interpretations, and manuscript preparation. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/falgy.2021.677019/full#supplementary-material
patients with severe and moderate atopic dermatitis. PLoS ONE. (2016) 11:e0156077. doi: 10.1371/journal.pone.0156077

28. Hedavati MT, Arabzadehmoghadam A, Hajheydari Z. Specific ige against alternaria alternata in atopic dermatitis and asthma patients. Eur Rev Med Pharmacol Sci. (2009) 13:187–91.

29. Salo PM, Yin M, Arbes SJ Jr, Cohn RD, Sever M, Burge HA, et al. Dustborne alternaria alternata antigens in us homes: results from the national survey of lead and allergens in housing. J Allergy Clin Immunol. (2005) 116:623–9. doi: 10.1016/j.jaci.2005.05.030

30. Zukiwczek-Sobczak WA. The role of fungi in allergic diseases. Postepy dermatologii i alergologii. (2013) 30:402–5. doi: 10.5114/pdia.2013.33377

31. Galand C, Leyva-Castillo WM, Yoon J, Han A, Lee MS, McKenzie ANJ, et al. Il-33 promotes food anaphylaxis in epicutaneously sensitized mice by targeting mast cells. J Allergy Clin Immunol. (2016) 138:1536–66. doi: 10.1016/j.jaci.2016.03.056

32. Peng G, Mu Z, Cui L, Liu P, Wang Y, Wu W, et al. Anti-il-33 antibody has a therapeutic effect in an atopic dermatitis murine model induced by 2, 4-dinitrochlorobenzene. Inflammation. (2018) 41:154–63. doi: 10.1007/s10753-017-0673-7

33. Scheffchick A, Kiriti D, Magin TM. Keratin defects influence the itch-inducing cytokine thymic stromal lymphopoietin through amphiregulin-epidermal growth factor receptor signaling. J Allergy Clin Immunol. (2019) 144:179–22. doi: 10.1016/j.jaci.2019.07.041

34. Bhagavatula N, Nerusu KC, Fisher GI, Liu G, Thakur AB, Gremmell L, et al. Amphiregulin and epidermal hyperplasia: amphiregulin is required to maintain the psoriatic phenotype of human skin grafts on severe combined immunodeficient mice. Am J Pathol. (2005) 166:1009–16. doi: 10.1016/S0002-9440(05)62322-X

35. Rojahn TB, Vorstandlechner V, Krausgruber T, Bauer WM, Alkon N, Bangert ANJ, et al. Il-33 antibody has a therapeutic effect in an atopic dermatitis murine model induced by 2, 4-dinitrochlorobenzene. Inflammation. (2018) 41:154–63. doi: 10.1007/s10753-017-0673-7

36. van den Oord RA, Sheikh A. Filaggrin gene defects and risk of developing allergic sensitisation and allergic disorders: systematic review and meta-analysis. BMJ. (2009) 339:b2433. doi: 10.1136/bmj.b2433

37. Leyva-Castillo JM, Hener P, Jiang H, Li M. Tspl produced by keratinocytes promotes allergen sensitization through skin and thereby triggers atopic march in mice. J Invest Dermatol. (2013) 133:154–163. doi: 10.1097/jid.2012.239

38. Banzt SK, Zhu Z, Zheng T. The atopic march: progression from atopic dermatitis to allergic rhinitis and asthma. J Clin Cell Immunol. (2014) 5:202. doi: 10.4172/2155-9899.1000202

39. Koppelman SJ, Knol EF, Vlooswijk RA, Wensing M, Knulst AC, Heffe SL, et al. Peanut allergen ara h 3: isolation from peanuts and biochemical characterization. Allergy. (2003) 58:1144–51. doi: 10.1034/j.1398-9999.2003.00259.x

40. Muto T, Fukuoka A, Kabashima K, Ziegler SF, Nakayoshi K, Matsushita K, et al. The role of basophils and proadergic cytokines, tslp and il-33, in cutaneously sensitized food allergy. Int Immunol. (2014) 26:539–49. doi: 10.1093/intimm/dxu058

41. Warrin S, Bernard H, Wal JM, Adel-Patient K. Cutaneous or respiratory exposures to peanut allergens in mice and their impacts on subsequent oral exposure. Int Arch Allergy Immunol. (2014) 161:189–99. doi: 10.1159/000363444

42. Monticelli LA, Osborne LC, Noti M, Tran SV, Zais DM, Arts D. Il-33 promotes an innate immune pathway of intestinal tissue protection dependent on amphiregulin-egf interactions. Proc Natl Acad Sci USA. (2015) 112:10762–7. doi: 10.1073/pnas.1509070112

43. Stoll SW, Stuart PE, Swindell WR, Tsoi LC, Li B, Gandarillas A, et al. The egf receptor ligand amphiregulin controls cell division via fomx1. Oncogene. (2016) 35:2075–86. doi: 10.1038/onc.2015.269

44. Pohin M, Guesdon W, Mekouo AA, Rabeony H, Paris I, Atanassov H, et al. Oncostatin m overexpression induces skin inflammation but is not required in the mouse model of imiquimod-induced psoriasis-like inflammation. Eur J Immunol. (2016) 46:1737–51. doi: 10.1002/eji.201546216

45. Stoll SW, Stuart PE, Lambert S, Gandarillas A, Rittie L, Johnston A, et al. Membrane-tethered intracellular domain of amphiregulin promotes keratinocyte proliferation. J Invest Dermatol. (2016) 136:444–52. doi: 10.1016/j.jid.2015.10.061

46. Yao H, Hao S, Andersen-Nissen E, Mauck WM, Zheng S, Butler A, et al. Integrated analysis of multimodal single-cell data. bioRxiv - Genom. (2020) 18:3537–87. doi: 10.1101/2020.12.333331

47. Franzen O, Gan L-M, Bjørgkjen JLM. Panglaodb: a web server for exploration of mouse and human single-cell rna sequencing data. Database. (2019) 2019:baz046. doi: 10.1093/database/baz046

48. Heng TSP, Painter MW, Elpek K, Lukacs-Kornek V, Mauermann N, Turley SJ, et al. The immunological genome project: networks of gene expression in immune cells. Nat Immunol. (2008) 9:1091–4. doi: 10.1038/nmi1091-1091

49. Valenzil E, Bulik M, Tabib T, Morse C, Sembrat J, Trejo Bittar H, et al. Single-cell analysis reveals fibroblast heterogeneity and myofibroblasts in systemic sclerosis-associated interstitial lung disease. Ann Rheumatic Dis. (2019) 78:1379–87. doi: 10.1136/annrheumdis-2018-214865

50. Joost S, Annuusker J, Jacob T, Sun X, Dalessandri T, Sivan U, et al. The molecular anatomy of mouse skin during hair growth and rest. Cell Stem Cell. (2020) 26:441–57. doi: 10.1016/j.stem.2020.01.012

51. Waterhouse J, Weinert D, Nevill A, Atkinson G, Reilly T. Some factors influencing the sensitivity of body temperature to activity in neonates. Pediatr Res. (2013) 30:42–5. doi: 10.1034/j.1398-9995.2013.00259.x

52. Alexander G. Body temperature control in mammalian young. J Perinatol. (2012) 32:16–24. doi: 10.1038/jp.2011.223

53. Buelow et al. Food Allergy Initiation: Alternaria Function
Cook-Mills JM, Kaplan MH, Turner MJ, Kloepfer KM, Kumar R. Correction: mechanism for initiation of food allergy: dependence on skin barrier mutations and environmental allergen costimulation. J Allergy Clin Immunol. (2018) 142:339.

Anvari S, Miller J, Yeh CY, Davis CM. Ige-mediated food allergy. Clin Allergy Immunol. (2019) 57:244–60. doi: 10.1007/s12016-018-8710-3

Kanagaratham C, El Ansari YS, Lewis OL, Oettgen HC. Ige and igg antibodies as regulators of mast cell and basophil functions in food allergy. Front Immunol. (2020) 11:603050. doi: 10.3389/fimmu.2020.603050

Saluja R, Khan M, Church MK, Maurer M. The role of il-33 and

Klein O, Sagi-Eisenberg R. Anaphylactic degranulation of mast

Espinosa E, Valitutti S. New roles and controls of mast cells: focus on compound exocytosis. J Immunol Res. (2019) 2019:9542656. doi: 10.1155/2019/9542656

Kubo M. Mast cells and basophils in allergic inflammation. Curr Opin Immunol. (2018) 54:74–79. doi: 10.1016/j.coi.2018.06.006

Bulfone-Paus S, Nilsson G, Draber P, Blank U, Levi-Schaffer F. Positive and negative signals in mast cell activation. Trends Immunol. (2017) 38:657–67. doi: 10.1016/j.it.2017.01.008

Espinosa E, Valitutti S. New roles and controls of mast cells. Curr Opin Immunol. (2018) 50:39–47. doi: 10.1016/j.coi.2017.10.012

Komi DEA, Mortaz E, Amani S, Tiotiu A, Folkerts G, Adcock IM. The role of mast cells in ige-independent lung diseases. Curr Rev Immunol. (2020) 58:377–87. doi: 10.1007/s12016-020-08779-5

Lyons DO, Pullen NA. Beyond ige: alternative mast cell activation across different disease states. Int J Mol Sci. (2020) 21:1498. doi: 10.3390/ijms21041498

Pemberton AD, Wright SH, Knight PA, Miller HR. Anaphylactic release of mucosal mast cell granule proteases: role of serpins in the differential clearance of mouse mast cell proteases-1 and –2. J Immunol. (2006) 176:899–904. doi: 10.4049/jimmunol.176.2.899

Snelgrove RJ, Gregory LG, Peiró T, Akhtar S, Campbell GA, Walker SA, et al. Alternaria-derived serine protease activity drives il-33-mediated asthma exacerbations. J Allergy Clin Immunol. (2014) 134:583–92. doi: 10.1016/j.jaci.2014.02.002

Richards CD. The enigmatic cytokine oncostatin m and roles in disease. ISRN Inflamm. (2013) 2013:521013. doi: 10.1155/2013/521013

Francis A, Bosio E, Stone SF, Fatovich DM, Arendts G, MacDonald SPJ, et al. Markers involved in innate immunity and neutrophil activation are elevated during acute human anaphylaxis: validation of a microarray study. J Innate Immun. (2019) 11:63–73. doi: 10.1159/000492301

Arshad MI, Guihard P, Danger Y, Noel G, Le Seyec J, Boutet MA, et al. Oncostatin m induces il-33 expression in liver endothelial cells in mice and expands st2+cd4+ lymphocytes. Am J Physiol Gastrointest Liver Physiol. (2015) 309:G542–53. doi: 10.1152/ajpgi.00398.2014

Richards CD, Izakelian L, Dubey A, Zhang G, Wong S, Kwofie K, et al. Regulation of il-33 by oncostatin m in mouse lung epithelial cells. Mediators Inflamm. (2016) 2016:9858374. doi: 10.1155/2016/9858374

Zeng F, Chen H, Chen L, Mao J, Cai S, Xiao Y, et al. An autocrine circuit of il-33 in keratinocytes is involved in the progression of psoriasis. J Invest Dermatol. (2020) 141:596–606. doi: 10.1016/j.jid.2020.07.027

Hayes T, Rumore A, Howard B, He X, Luo M, Wuenschmann S, et al. Innate immunity induced by the major allergen alt a 1 from the fungus alternaria is dependent upon toll-like receptors 2/4 in human lung epithelial cells. Front Immunol. (2018) 9:1507. doi: 10.3389/fimmu.2018.01507

Jiao D, Wong CK, Qiu HN, Dong J, Cai Z, Chu M, et al. Nod2 and tlr2 ligands trigger the activation of basophils and eosinophils by interacting with dermal fibroblasts in atopic dermatitis-like skin inflammation. Cell Mol Immunol. (2016) 13:535–50. doi: 10.1038/cmi.2015.77

Svanaprakorn P, Tongyen T, Prakhongcheep O, Laoratthaphong P, Chanvorachote P. Establishment of an anti-acne vulgaris evaluation method based on tlr2 and tlr4-mediated interleukin-8 production. In Vivo. (2019) 33:1929–34. doi: 10.21873/invivo.11687

Chinthrajah S, Cao S, Liu C, Lye SC, Sindher SB, Long A, et al. Phase 2a randomized, placebo-controlled study of anti-il-33 in peanut allergy. JCI insight. (2019) 4:e131347. doi: 10.1172/jci.insight.131347

Brandt EB, Sivaprasad U. Th2 cytokines and atopic dermatitis. J Clin Cell Immunol. (2011) 2:110. doi: 10.4172/2155-9899.1000110

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