Flightless I Alters the Inflammatory Response and Autoantibody Profile in an OVA-Induced Atopic Dermatitis Skin-Like Disease

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Atopic dermatitis (AD) is a chronic pruritic inflammatory skin disease characterized by excessive inflammation and disrupted skin barrier function. Although the etiology of AD is not completely understood, clinical and basic studies suggest increasing involvement of autoantibodies against intracellular proteins. An actin remodeling protein, Flightless I (Flii), has been shown to promote development of inflammatory mediated skin conditions and impairment of skin barrier development and function. Here, we sought to determine the effect of altering Flii expression on the development of AD and its contribution to autoimmune aspects of inflammatory skin conditions. Ovalbumin (OVA)-induced AD skin-like disease was induced in Flii heterozygous (Flii+/−), wild-type (Flii+/+), and Flii transgenic (Flii Tg/Tg) mice by epicutaneous exposure to OVA for 3 weeks; each week was separated by 2-week resting period. Reduced Flii expression resulted in decreased disease severity and tissue inflammation as determined by histology, lymphocytic, and mast cell infiltrate and increased anti-inflammatory IL-10 cytokine levels and a marked IFN-γ Th1 response. In contrast, Flii over-expression lead to a Th2 skewed response characterized by increased pro-inflammatory TNF-α cytokine production, Th2 chemokine levels, and Th2 cell numbers. Sera from OVA-induced AD skin-like disease Flii+/− mice showed a decreased level of autoreactivity while sera from Flii Tg/Tg mice counterparts showed an altered autoantibody profile with strong nuclear localization favoring development of a more severe disease. These findings demonstrate autoimmune responses in this model of OVA-induced AD-like skin disease and suggest that Flii is a novel target, whose manipulation could be a potential approach for the treatment of patients with AD.

Keywords: atopic dermatitis, flightless I, autoantibody, inflammation, skin barrier

INTRODUCTION

Atopic dermatitis (AD) is one of the most common heterogeneous inflammatory skin diseases affecting 20% of children and 1–3% of adults worldwide (1). The disease is associated with impairments in the skin barrier and variable clinical indicators including occurrence of eczematous lesions, pruritus, and cheilitis (1). The etiology of AD is complex and is often characterized by abnormal immunological pathways that manifest in an imbalance of T-helper (Th)1 and Th2 responses (2). Typically, AD is described as having a biphasic course consisting of an acute inflammatory Th2-dominated phase associated with IgE production and a chronic phase distinguished by reappearance...
of Th1 responses, tissue remodeling, and dermal thickening (3). Histopathologically, AD is characterized by an inflammatory infiltrate consisting of CD4+ memory T cells, mast cells, and eosinophils and a controlled temporal–spatial expression of pro-inflammatory cytokines and chemokines driving atopic inflammation of the skin (4).

Research within the last decade has found an association between AD and autoantibody development, suggesting the contribution of autoantibodies to the pathogenesis of AD (3, 5–8). It is proposed that tissue damage induced by AD allows exposure of intracellular antigens that are normally inaccessible by antibodies to the extracellular space, where they can interact with B cells and antibodies (9). A broad spectrum of IgE targeting self-antigens have been identified in above 90% of severe AD patients and high-avidity IgG autoantibodies have been proposed as potential diagnostic markers for severe AD (5, 10–12). The fact that these autoantibodies are associated with disease severity implicates their role in both humoral and cellular immunity in AD pathogenesis (5, 12). Antinuclear antibodies have been shown to have both biological and clinical significance acting as sensors of cellular stress and inflammation associated with environmental factors (13). While the presence of autoantibodies can have a protective role as natural autoantibodies (14), the presence of nuclear autoantibodies in AD has been suggested to lead to the continual provocation of the immune system hence contributing to the severity and chronicity of disease.

Flightless I (Flii) is a highly conserved and unique member of the gelsolin family of actin remodeling proteins and a nuclear receptor activator affecting the transcriptional activity of many modulators of tissue remodeling and inflammation (15, 16). Flii has been demonstrated to regulate cytokine secretion and cellular inflammatory responses via its intracellular and extracellular effects on toll-like receptors (17–20). Flii expression increases in skin during development and in response to inflammation, injury, skin cancer development, wound healing, and skin blistering (21–23). Over-expression of Flii delays the development of an intact skin barrier in the embryonic skin of mice and impairs the recovery of the epidermal barrier post injury via its effects on tight-junction formation (22). A recent study has shown that reducing Flii levels either genetically or using Flii neutralizing antibodies decreases erythema, inflammatory cell infiltrate, and pro-inflammatory cytokine secretion in a mouse model of psoriasisform dermatitis (24). Taken together, these findings suggest a possible role for Flii in inflammatory responses mediating AD. Using an OVA-induced AD-like skin mouse model (25), this study aimed to investigate the effect of differential Flii gene expression on development of AD via characterization of inflammatory and autoimmune responses responsible for AD pathogenesis.

**MATERIALS AND METHODS**

**Animal Studies**

Mice were maintained according to the Australian Code for the Care and Use of Animals for Scientific Purposes under protocols approved by the Child Youth and Women’s Health Service Animal Ethics Committee (AEC916/06/2015). Mice with the BALB/c background and wild-type controls were obtained from inbred litters. Flii-deficient heterozygous null mice (Flii+/−) and mice carrying the complete human Flii gene on a cosmids transgene were maintained as described previously (26, 27). Heterozygous transgenic mice Flii+/− were made by crossing Flii+/− with cosmids transgene Flii+/+. These transgenic mice were intercrossed to obtain animals homozygous for the transgene Flii+/+ which carry two copies of the mouse Flii gene and two copies of the human Flii transgene.

Atopic dermatitis was induced via epicutaneous exposure of mice to OVA as previously described (25). Briefly, 10- to 12-week-old female wild-type (Flii+/-), Flii heterozygous (Flii+/−), and Flii transgenic (Flii+/-/+) mice (n = 8/genotype) were anesthetized using isoflurane inhalation, the skin on the back of mice was shaved and then tape stripped four times by a transparent adhesive tape (Tegaderm) to introduce a standardized skin injury. A gauze patch (1 x 1 cm²) soaked with 100 µl of 0.1% OVA (OVA group) in saline or 0.9% saline (control group) was placed on the back skin and secured with Tegaderm dressing. The experiment comprised three 1-week exposures with a 2-week interval between each exposure week (Figure 1A). Clinical images of affected skin, transepidermal water loss (TEWL), and erythema measurements were recorded at the end of the third sensitization week. The level of skin erythema was measured daily using a handheld DermaLab Unit (Cortex Technology) following manufacturer’s instructions. This instrument uses skin reflectance spectroscopy to determine the redness of inflamed skin. The instrument was blanked prior to placing a probe directly onto the inflamed AD-like lesions on dorsal skin and a reading obtained as previously described (24). Measurements of TEWL were obtained with a calibrated Vapometer evaporimeter (Delfin Technologies, Finland). The probe was allowed to equilibrate for approximately 30 s before a brief measurement period (8 s) as indicated in manufacturer’s guidelines. The Vapometer was placed directly onto the inflamed AD-like lesions on dorsal skin (using an 11-mm adaptor) and held there securely for the duration of the measurement. For both erythema and TEWL analysis, three separate measurements were taken per mouse to ensure the entire region of AD-like inflamed back skin was assessed as previously described (24). Mice were then euthanized and skin biopsies collected for histology analysis, immunohistochemistry, and mRNA extraction. Blood was collected for autoimmune experiments.

**Histology and Immunohistochemistry**

Paraffin-embedded fixed tissue samples were stained with hematoxylin and eosin or toluidine blue following established protocols (28). Level of skin inflammation was assessed using skin thickness measurements, and analysis of inflammatory and mast cell numbers in lesional skin of OV A-induced AD-like skin lesions (24). Measurements of TEWL were obtained with a calibrated Vapometer evaporimeter (Delfin Technologies, Finland). The probe was allowed to equilibrate for approximately 30 s before a brief measurement period (8 s) as indicated in manufacturer’s guidelines. The Vapometer was placed directly onto the inflamed AD-like lesions on dorsal skin (using an 11-mm adaptor) and held there securely for the duration of the measurement. For both erythema and TEWL analysis, three separate measurements were taken per mouse to ensure the entire region of AD-like inflamed back skin was assessed as previously described (24). Mice were then euthanized and skin biopsies collected for histology analysis, immunohistochemistry, and mRNA extraction. Blood was collected for autoimmune experiments.
mast cells in entire sections of OVA-induced AD-like skin lesions per mm². Immunohistochemistry was also performed on paraffin-embedded fixed AD-like skin lesions following antigen retrieval according to the manufacturer’s protocols (DAKO Corporation, Glostrup, Denmark). Following blocking in 3% normal goat serum, primary antibodies against CD4 [rat monoclonal, #14-9766-82 (Thermo Scientific, Australia) (1:200)], T-bet [rabbit polyclonal, #PA5-40573 (Thermo Scientific, Australia) (1:100)], GATA-3 [rabbit polyclonal, #ab106625 (Abcam Australia) (1:100)], and ROR-γ [rabbit monoclonal, #ab207082 (Abcam Australia) (1:3,000)] were applied and slides were incubated at 4°C overnight in a humidified chamber before application of species-specific, Alexa Flour-488 or Alexa Flour-594 secondary antibodies (Invitrogen, Australia) for 1 h at room temperature. Finally, slides were washed and mounted in Fluorescence Mounting Medium (Dako, Australia). Images were captured on Olympus microscope and CellSense Live Science Imaging Software program (Olympus, Germany) used for counting the positive cells in the AD-like lesions of Flii+/−, wild-type, and FliiTg/Tg mice. Negative controls included replacing primary antibodies with normal rabbit IgG, or normal mouse IgG. For verification of staining, non-specific binding was determined by omitting primary or secondary antibodies. All control sections had negligible immunofluorescence.

**RTq-PCR**

Harvested tissue was snap-frozen in liquid nitrogen and total RNA was isolated using Ultraclean Tissue and Cell RNA Isolation
Kit (MoBio Laboratories, Carlsbad, CA, USA) according to the manufacturer’s protocol. Total cDNA was synthesized using iScript cDNA Synthesis Kit (Bio-Rad Laboratories, Hercules, CA, USA) according to manufacturer’s protocol. Quantitative PCR was performed using iQ SYBR Green Supermix (Bio-Rad Laboratories, Hercules, CA, USA) in triplicate reactions. The plates were placed in a CFX Connect Real-Time PCR Detection System (Bio-Rad Laboratories, Hercules, CA, USA). Reactions underwent 30 s at 95°C, then 40 cycles of 5 s at 95°C, 20 s at 60°C, and 10 s at 95°C before determination of melt curve between 65 and 95°C. GAPDH and CyPA were used as reference genes and inter-reaction calculator method was applied for all plates. For relative comparison, the cycle threshold value (Ct) was analyzed using the ΔΔCt method and data reported as Ct normalized to reference genes. Sequences for PCR primers are listed in Table S1 in Supplementary Material.

Autoantibody Immunofluorescence

In order to assess the degree of autoimmunity in OVA-induced AD-like skin of Flii+/−, wild-type, and Flii+/+− mice, sub-confluent primary wild-type mouse keratinocytes were stained with mouse sera of AD-induced mice following established protocols (13, 29). Briefly, primary keratinocytes were isolated from murine epidermis as previously described (30), grown on glass coverslips and washed in 1 × phosphate-buffered saline before paraformaldehyde (4%) fixation (10 min at room temperature). Fixed cells were subsequently permeabilized using 0.2% Triton-X-100 and 0.5% BSA in 1 × phosphate-buffered saline (5 min at room temperature) before incubation with pooled murine serum (n = 3) diluted (1 in 10) in 0.5% BSA in 1 × phosphate-buffered saline for 1 h. Bound murine IgG was detected with Alexa Flour 633 goat anti-mouse IgG (1 in 1,000; #A21050; Invitrogen, Mulgrave, VIC, Australia) in 1 × phosphate-buffered saline for 1 h at room temperature. Following repeated 2 min washes with 1 × phosphate-buffered saline, cells were stained with DAPI nucleic acid stain (Sigma-Aldrich) for 5 min at room temperature before washing and mounting for imaging. Negative controls included omitting the incubation with mouse serum. All control sections had negligible immunofluorescence. Staining pattern of autoantibodies was assessed using the Olympus microscope and CellSense Live Science Imaging Software program (Olympus, Germany).

Immunoblot Analysis of AD Serum Autoreactive Antibodies

Protein was extracted from WT murine keratinocytes using standard protein extraction protocols (31). Samples of extracted protein (20 µg) were run on 10% SDS-PAGE gels (30 min; 200V) and transferred to nitrocellulose by wet transfer (1 h; 100V). Membranes were cut into strips, blocked in 3% bovine serum albumin (Sigma) for 30 min, and probed with murine serum diluted (1 in 10) in tris-buffered saline containing 3% BSA and 0.1% Tween overnight. After washing, horseradish peroxidase-conjugated goat anti-murine immunoglobulin secondary antibody was added for a further 1 h at room temperature. Stringent washes were performed before detection of horseradish peroxidase and exposure using GeneSnap analysis program (SynGene, Frederick, MD, USA).

Statistical Analysis

Statistical differences were determined using the Student’s t-test or one-way ANOVA. For data not following a normal distribution, the Mann–Whitney U test was performed. A P value of less than 0.05 was considered significant.

RESULTS

Flii-Deficient Mice Exhibit Reduced OVA-Induced AD-Like Skin Disease

The repeated epicutaneous exposure of OVA (Figure 1A) on mouse skin results in an AD-like skin disease with histological and phenotypic features of human AD, including erythema, skin thickening, and a localized immune response (25). Flii homozygous (Flii+/−) mice are embryonic lethal (26); therefore, using the OVA-induced AD-like skin mouse model, the severity of OVA-induced AD-like lesions was determined in response to different Flii gene levels in Flii heterozygous (Flii+/−), normal (Flii+/+), and Flii transgenic (Flii+/+−) mice. All three mice genotypes developed localized inflammation which increased following each week of OVA exposure. Flii-deficient mice showed reduced levels of inflammation and scaling (Figure 1B). Spectrophotometric measurement of the redness of the OVA exposed skin showed that Flii-deficient mice had significantly less erythema (Figure 1B) than wild-type and Flii+/+− counterparts at day 50 of the experiment. Similarly, the degree of TEWL was also significantly reduced in Flii+/− mice and very similar to that observed in control animals (Figure 1C). Control mice administered saline only showed no evidence of AD-like inflammation macroscopically or any development of erythema or TEWL (Figures 1B–D).

A hallmark of AD is skin thickening with a marked influx of leukocyte and mast cell inflammatory infiltrate. All OVA-induced AD-like skin sections showed evidence of AD compared to control saline exposed skin including a degree of epidermal hyperplasia and edema coupled with inflammatory lymphocytic and mast cell dermal infiltrate (Figures 2A–C). Examining the skin thickness in OVA-induced AD-like skin lesions revealed that deficient Flii mice had significantly thinner skin than OVA-induced AD-like skin lesions of wild-type and Flii+/+− mice (Figure 2A). Assessment of lymphocytic and mast cell dermal infiltrate showed that Flii deficiency consistently resulted in significantly decreased inflammatory cell numbers (Figures 2B,C). In contrast, both normal and increased Flii gene expression resulted in thickened epidermis, increased numbers of inflammatory cells (Figure 2B) and mast cells (Figure 2C). Control mice skin exposed to saline only gave patch showed no evidence of AD-like dermatitis features macroscopically and had low levels of inflammatory infiltrate in the dermis (Figures 2A–C).

Anti-Inflammatory Cytokine mRNA Profiles Are Increased in OVA-Induced AD-Like Skin of Flii-Deficient Mice

OVA-induced AD-like skin lesions of Flii+/−, wild-type, and Flii+/+− mice were assessed for Flii and levels of cytokines and chemokines mediating the Th1 and Th2 inflammatory responses.
during AD pathogenesis. Flii-deficient mice showed approximately 25% decrease in Flii levels, while Flii over-expressing mice had a twofold increase in Flii levels compared to wild-type counterparts (Figure 3). Flii gene levels were found to affect number of key cytokines and chemokines responsible for development of AD-like lesions. Notably, main Flii deficiency favored a Th1 immune response and decreased inflammation with significant threefold increase in IFN-γ mRNA levels compared to FliiTg/Tg mice, decreased pro-inflammatory IL-4, and increased anti-inflammatory IL-10 mRNA levels compared to both wild-type and FliiTg/Tg mice (Figure 3). Interestingly, Flii−/- OVA-induced AD-like skin lesions also showed increased IL-5 and IL-6 mRNA levels when compared to FliiTg/Tg mice but not wild-type counterparts (Figure 3). In contrast, over-expression of Flii resulted in similar cytokine levels to wild-type counterparts, except significantly increased TNF-α mRNA expression and significantly reduced IFN-γ mRNA expression when compared to both Flii-deficient and wild-type mice, which would favor more severe AD manifestation. Cytokine mRNA levels of IL-13, IL-23, and IL-17A did not differ between OVA-induced AD-like skin lesions of three genotypes (Figure 3). Expression of CCL22 mRNA was significantly increased in OVA-induced AD-like skin lesions of FliiTg/Tg mice compared to wild-type mice, while OVA-induced AD-like skin lesions of Flii−/- mice showed elevated CCL17 chemokine mRNA expression compared to FliiTg/Tg mice (Figure 4). CXCL9 chemokine mRNA levels were found to be similar in all three genotypes (Figure 4). The unbalanced skewed ratio of Th1/Th2 response in FliiTg/Tg mice was confirmed by immunohistochemical co-staining of T cell subset markers, namely Th1 (CD4 and T-bet), Th2 (CD4 and GATA-3), and Th17 (CD4 and ROR-γ) showing increased Th2 cell numbers in dermal area of FliiTg/Tg mice (Figures 5A–D).

**FliiTg/Tg OVA-Induced AD-Like Disease Mice Have Altered Autoreactivity Profiles Compared With WT and Flii−/- Mice**

To compare autoantibody formation in the OVA-induced AD-like disease model between Flii-deficient, wild-type, and Flii-overexpressing mice, serum samples collected at day 50...
were assessed by immunofluorescence and immunoblot analysis. Primary keratinocytes from wild-type mice were fixed and probed with pooled sera from experimental mice and staining patterns analyzed using fluorescence microscopy. Cytoplasmic and perinuclear staining within keratinocytes was apparent after incubation with sera from all three genotypes, however, only sera from OVA-induced AD-like disease Flii<sup>Tg</sup> mice produced strong nuclear staining patterns (Figure 6A). To further elucidate autoreactivity in these samples, keratinocyte whole-cell lysates were separated via SDS-PAGE, transferred to nitrocellulose membrane, and probed with pooled sera (Figure 6B). Additionally, autoreactivity patterns of individual mouse sera are shown in Figure S1 in Supplementary Material. Regions of differential autoreactivity were detected at 145, 70, 60, and 43 kDa band sizes (black arrows). Immunoblot analysis of sera (Figure 6C) from individual mice found that autoreactivity was highest in wild-type mice (50%) and Flii<sup>Tg</sup> mice (43%), while Flii-deficient (Flii<sup>−/−</sup>) mice showed the lowest degree of autoreactivity (25%). Positive autoreactive bands at 70 kDa were also different in Flii<sup>Tg</sup> mice compared to the two other genotypes, which may be responsible for the nuclear staining pattern observed in immunofluorescence analysis. Immunoblot analysis of sera from normal non-dermatitis mice from each genotype were also analyzed; no autoreactivity was detected (Figure S2 in Supplementary Material).

**DISCUSSION**

Flii is a cytoskeletal protein with important roles in skin development, tight junction function, skin barrier establishment, and recovery post injury which is upregulated in response to tissue...
inflammation and wounding (22, 31). The secreted form of Flii has been shown to affect innate immune signaling pathways and modulate cell activity and cytokine secretion from fibroblasts and macrophages in vitro (17, 20). Recent studies have described the contribution of Flii over-expression in exacerbation of inflammatory conditions, including psoriasiform dermatitis (24) and an autoimmune inflammation-mediated epidermolysis bullosa acquisita (23, 32).

To determine if Flii affects AD development and severity, mice with low (Flii<sup>+/−</sup>), normal (Flii<sup>+/+</sup>), and high (FliiTg/Tg) expression of the Flii gene were exposed to repeated epicutaneous exposure to OVA and induced to form AD-like skin lesions. In contrast to our original hypothesis, Flii over-expression did not result in a more exacerbated development of erythema and AD-like skin lesions. However, reduced Flii levels in Flii<sup>−/−</sup> mice led to significantly decreased development of AD-like skin lesions as marked by decreased erythema, TEWL, epidermal hyperplasia, and lymphocytic and mast cell tissue infiltrate compared to both wild-type and FliiTg/Tg mice. Despite increased IFN-γ levels, the AD lesions of Flii<sup>+/−</sup> mice had reduced IL-4, which may contribute significantly to the reduced skin thickening in this model of AD (33). These findings are in agreement with studies which demonstrated that reducing Flii expression, either genetically or using Flii neutralizing antibodies, decreased tissue inflammation, and disease severity in mouse models of psoriasiform dermatitis and epidermolysis bullosa acquisita (23, 24, 32).

Genetically modified mice engineered to over-express Th2 cytokines develop skin barrier defects and AD spontaneously (34). The formation of AD lesions is known to be triggered by production of Th2 cytokines by mast cells and CD4<sup>+</sup> T cells which also promote IgE production by B cells, while Th1 cells secrete IFN-γ to suppress proliferation of Th2 cells and IgE synthesis (35, 36). The dominance of Th2 cytokines in AD cause decreased expression of filagrin and other barrier promoting molecules found in the skin (3). Examining the levels of cytokines and chemokines produced in AD-like lesions of Flii<sup>+/−</sup>, wild-type, and FliiTg/Tg mice revealed decreased IFN-γ levels in FliiTg/Tg mice suggestive of Th2 immune responses, while Flii<sup>+/−</sup> had increased IFN-γ levels and decreased clinical AD severity suggestive of potential Flii effect on IgE synthesis, however, this is yet to be investigated. Analysis of T-helper subsets within lesional skin showed a significantly Th2-skewed response in OVA-induced AD-like lesions of FliiTg/Tg mice with higher numbers of CD4<sup>+</sup>GATA3+ cells compared to AD-like lesions of Flii<sup>+/−</sup> or wild-type counterparts who showed significantly higher Th1:Th2 ratio. Additionally, AD-like lesions of FliiTg/Tg mice showed increased pro-inflammatory TNF-α cytokine levels and while TNF-α has been demonstrated to promote the AD development (37) we did not observe increased
AD severity in these mice in this model of OVA-induced AD. As mast cell numbers were not significantly increased in Fliitg/− mice, and T-helper cells were heavily skewed toward a Th2 phenotype within Fliitg/− lesions, excess TNF-α was likely produced by other cell types such as macrophages, which is further supported by increased CCL22 chemokine levels observed in Fliitg/− mice. Interestingly, previous studies have shown that secreted Flii reduces macrophage secretion of TNF-α in vitro (20) while in vivo studies using a mouse model of psoriasiform dermatitis showed reduced TNF-α levels in response to reduced Flii (24) which was not observed in this model of OVA-induced AD. These findings may reflect inherent differences between in vivo and in vitro studies, for example differences in complex in vivo, multicellular environments, as well as differences in different models of inflammatory skin diseases.

In agreement with previous studies demonstrating decreased tissue inflammation and inflammatory cytokine secretion in Flii-deficient mice (24, 32), OVA-induced AD-like skin lesions of Flii−/− mice showed a reduced inflammatory response marked by significantly increased anti-inflammatory IL-10 secretion as well as significantly increased IFN-γ and significantly reduced IL-4 levels. Indeed, IL-4 has been shown to be essential for eosinophil recruitment, Th2 cell differentiation, and IgE production (38). The increased CCL17 expression observed in AD lesions of Flii−/− mice compared to Fliitg/− mice may be a potential mechanism to restore Th1/Th2 balance, as CCL17 has previously been shown to induce a Th2-dominated inflammatory reaction (39). Reduction of Flii levels did not alter IL-23 and IL-17A cytokine levels as previously observed in psoriasiform dermatitis (24) suggesting that Th17 responses are not involved in this model of AD. Despite significant differences in IFN-γ levels between three genotypes, CXCL9 chemokine levels were not altered between genotypes in this model of AD. IL-13 levels were also not significantly different between three genotypes, however, there was a trend to higher levels in Fliitg/− mice OVA-induced AD-like skin lesions and IL-13 cytokine has previously been linked to autoantibody production in early rheumatoid arthritis (40).

Autoimmunity has been increasingly recognized to play part in exacerbating the severity of AD (6, 9, 12) as a consequence of both humoral and cellular immunity (6). It is postulated that IFN-γ signaling during AD pathogenesis may promote the development of autoimmunity as IFN-γ overexpressing mice spontaneously develop autoantibodies (41, 42) and deletion of the IFN-γ receptor inhibits autoantibody production in lupus-prone mice (43). On the basis of our findings showing altered AD severity and altered Th1/Th2 responses including significantly altered IFN-γ expression with different Flii genotypes, we examined the effect of Flii levels on autoimmunity in OVA-induced
AD-like skin lesions. The decreased severity of AD observed in *Flii*−/− mice also correlated with the reduced degree of autoactivity (50% reduction vs WT), and further studies are required to investigate whether the autoantibodies from OVA induced AD-like skin mice contribute to or are a product of the observed AD-like symptoms. Immunoblot analysis of sera from normal non-dermatitis mice showed no autoactivity suggesting that autoantibody development was disease-specific. Interestingly, both *Flii*−/− and wild-type mice showed similar autoantibody staining patterns and regions of differential autoactivity. *Flii*−/− mice showed similar levels of autoactivity compared to wild-type mice, however, in addition to the cytoplasmic and perinuclear staining pattern observed in all genotypes, the sera of these mice had a strong nuclear staining pattern. This pattern may indicate the presence of disease-mediating antinuclear antibodies which have been clinically associated with AD and other inflammatory skin disorders (44–46). Flii has previously been shown to affect TLR signaling pathways, both intracellularly and extracellularly, hence modulating innate inflammatory responses and directly impacting immune signaling, however, the potential role of Flii in autoimmunity has not been explored to date. Further studies are required to determine if autoactivity in *Flii* genetic mice is a direct result of the impact of Flii on immune signaling or if it is secondary to increased pathology observed in these animals. In addition, determining the subtype of autoreactive immunoglobulins developed in the OVA-challenged murine model of AD would allow more comparison to be drawn to findings of previous clinical cohorts (10, 47).

While major differences between human AD and murine models have been demonstrated (47, 48), models of AD promote our understanding of the complex pathogenesis of human AD, and identify potential novel targets for design of targeted biologics (49). Here, we have demonstrated that Flii is a novel target in AD and that reducing its levels decreased the severity of AD in the ovalbumin-challenged murine model of AD. Additionally, we have examined the role of autoimmunity in this model of AD and while the exact mechanisms are yet to be identified, our results suggest that the effects of Flii upon Th1/Th2 balance and autoimmunity are important during AD pathogenesis.

**ETHICS STATEMENT**

Mice were maintained according to the Australian Code for the Care and Use of Animals for Scientific Purposes under protocols approved by the Child Youth and Women’s Health Service Animal Ethics Committee (AEC916/06/2015).

**AUTHOR CONTRIBUTIONS**

AC and ZK conceived all the experiments with assistance from NS. ZK and NS carried out experiments and analysis with the assistance of HC and GY. ZK, NS, and AC wrote the manuscript and all authors contributed to the manuscript preparation and approved the final submitted and published versions.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at https://www.frontiersin.org/articles/10.3389/fimmu.2018.01833/full#supplementary-material.

**TABLE S1** | Primer sequences used in real-time qPCR.


