Calcineurin Aα Contributes to IgE-Dependent Mast-Cell Mediator Secretion in Allergic Inflammation

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\section*{Abstract}
Mast cells (MCs) are key mediators of allergic inflammation through the activation of cross-linked immunoglobulin E (IgE) bound to the high-affinity IgE receptor (FceRI) on the cell surface, leading to the release of biologically potent mediators, either from preformed granules or newly synthesized. Pharmacological inhibitors have been developed to target a key signaling protein phosphatase in this pathway, calcineurin, yet there is a lack of genetic and definitive evidence for the various isoforms of calcineurin subunits in FceRI-mediated responses. In this study, we hypothesized that deficiency in the calcineurin Aα isoform will result in a decreased allergic immune response by the MCs. In a model of passive cutaneous anaphylaxis, there was a reduction in vascular permeability in MC-deficient mouse tissues reconstituted with calcineurin subunit A (CnAα) gene-knockout (CnAα\textsuperscript{−/−}) MCs, and in vitro experiments identified a significant reduction in release of preformed mediators from granules. Furthermore, released levels of de novo synthesized cytokines were reduced upon FceRI activation of CnAα\textsuperscript{−/−} MCs in vitro. Characterizing the mechanisms associated with this deficit response, we found a significant impairment of nuclear factor of kappa light polypeptide gene enhancer in B cell phosphorylation and impaired nuclear factor kappa-light-chain-enhancer of activated B-cell inhibitor alpha (NF-κB) activation. Thus, we concluded that CnAα contributes to the release of preformed mediators and newly synthesized mediators from FceRI-mediated activation of MCs, and this regulation includes NF-κB signaling.

\section*{Introduction}
Allergic inflammation, or allergy, is an excessive and inappropriate immune response against specific though otherwise harmless allergens that the immune system had been previously sensitized to [1]. The inflammation in the allergic response is a result of a coordination of signaling cascades of various immune cell types and secretory mechanisms [2]. One immune cell type identified as the conductor in allergy and ensuing pathology is the mast cell (MC). Activation of the MC by an allergen elicits the
release of a plethora of inflammatory products that have effects manifesting in the overall allergic inflammatory response [3–5]. As MCs are tissue-resident sentinel cells populating sites of host-external environment interfaces, they are primed by specific immunoglobulin E (IgE) following the initial allergen exposure and subsequently coordinate the response through the release of potent mediators on a challenge exposure. Thus, it is crucial to understand the mechanisms underlying MC activation by the allergen.

MC-allergen interactions are mediated by IgE bound to the tetrameric high-affinity IgE receptor, FcεRI. FcεRI comprises 3 subunits, an α-chain that binds to IgE, a β-chain that amplifies signaling, and 2 γ-chains that are responsible for signal transduction through immunoreceptor tyrosine-based activation motifs. Aggregation and cross-linking of antigen (Ag)-bound IgE on FcεRI result in the internalization of the complex and initiation of the FcεRI-mediated signaling pathway. The FcεRI-mediated signaling pathway, recently reviewed by others, is highly complex [6, 7]. Briefly, multiple signaling pathways are activated including nuclear factor of activated T cells, mitogen-activated protein kinases (MAPKs), and nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) [8]. The FcεRI signaling cascade essentially results in 2 main events: (1) degranulation of preformed mediators and (2) de novo production and secretion of lipid mediators, cytokines, and chemokines.

Calcineurin is a serine-threonine protein phosphatase conserved in all eukaryotes with important functions in multiple cell types and signaling pathways [9]. Calcineurin is composed of a catalytic A (CnA) and regulatory B (CnB) subunit, and each subunit contains specific domains that promote the activation of calcineurin in a calcium-dependent manner. The catalytic subunit consists of 3 isoforms – α (CnAα), β (CnAβ), and γ (CnAγ). CnAα and CnAβ are ubiquitously expressed, while CnAγ is found in limited tissues [10]. Binding of calcium to CnB and calmodulin promotes conformational changes to calcineurin and the removal of an auto-inhibitory domain on CnA, allowing interactions with substrates [11–13]. The interaction between calcineurin and a variety of substrates regulates many physiological functions, ranging from cell cycle and apoptosis to immune responses [14, 15]. We reported that the regulator of calcineurin (Rcan) negatively regulates FcεRI-mediated allergic inflammation so we are still lacking genetic and definitive evidence of the mechanistic roles that calcineurin plays in the FcεRI-mediated immune response to an allergen – specifically the contributions of the different isoforms.

There has been limited research implicating CnAα in physiological settings. The homozygous CnAα gene deficiency results in early lethality in mice pups, a limitation to studying gene knockout mice [23]. One group overcame this obstacle by dietary intervention to show CnAα deficiency impaired in vivo T-cell responses to the antigen [24]. Otherwise, there is a paucity of literature or research on this calcineurin isoform in MCs and the FcεRI-mediated immune response despite the use of calcineurin inhibitors to treat allergy. Thus, it is important to delineate and understand the mechanisms in which CnAα contributes to FcεRI-mediated allergic inflammation so that improvements can be made in therapeutic approaches concerning calcineurin. In this study, using cells cultured and differentiated from homozygous knockout pups’ livers, we identified CnAα as a contributor to the early phase of FcεRI-mediated MC-specific allergic inflammation events.

**Materials and Methods**

**Animals**

CnAα−/− mice were generously provided by Dr. Jennifer Gooch (Emory University, Atlanta, GA, USA). MC-deficient KitW−/− mice were purchased from the Jackson Laboratory, Bar Harbor, ME, USA (B6Cg-kit W-sh/HNihAcBsmJ NistltF4). Mice colonies were purchased from the Jackson Laboratory, Bar Harbor, ME, USA in accordance with the guidelines of the Canadian Council of Animal Care.

**Antibodies and Reagents**

The anti-calcineurin Aα antibody was purchased from Millipore (Billerica, MA, USA). Antibodies to phospho-p38 mitogen-activated protein kinase (MAPK) (Thr 180/Tyr 182), phospho-JNK (Thr 183/Tyr 185), JNK, phospho-ERK1/2 (Thr202/Tyr204), factor kappa-light-chain-enhancer of activated B cells and p65 (RelA) (Ser 536) were generously provided by Dr. Jennifer Gooch (Emory University, Atlanta, GA, USA). MC-deficient KitW−/− mice were purchased from the Jackson Laboratory, Bar Harbor, ME, USA (B6Cg-kit W-sh/HNihAcBsmJ NistltF4). Mice colonies were purchased from the Jackson Laboratory, Bar Harbor, ME, USA (B6Cg-kit W-sh/HNihAcBsmJ NistltF4). Mice colonies were purchased from the Jackson Laboratory, Bar Harbor, ME, USA (B6Cg-kit W-sh/HNihAcBsmJ NistltF4).
MC Culture and Activation

Mouse liver-derived MCs (LMCs) were obtained by culturing liver cells from neonatal mice from CnAα+/− parents. Briefly, liver tissue was removed to a laminar flow hood and ground to produce a cell suspension in Roswell Park Memorial Institute (RPMI) 1640 medium, passed through a 40-μm cell strainer, then centrifuged at 500 g for 5 min at 4°C and finally resuspended at a density of 0.5 × 10^6 cells/mL in complete medium. Complete MC media consists of RPMI 1640 with 1-glutamine, 10% heat-inactivated fetal bovine serum (Gibco, Thermo Fisher Scientific), 4 μL SsoAdvanced Universal SYBR Green Supermix (Bio-Rad Laboratories), and data processed on Bio-Rad CFX Manager 3.1 software. Melt curve analyses were run to determine target specificity. Gene expression levels were normalized to hypoxanthine guanine phosphoribosyltransferase mRNA and analyzed using the ΔΔCt method. Primer sequences are reported in Table 1.

Flow Cytometry, Degranulation, and Calcium Mobilization

Samples from monocytes were stained directly using either FITC-CD117, FITC-IgG2b (FITC-CD117 isotype control), FITC-IgE, or FITC-rat IgG1 (FITC-IgE isotype control) antibodies and the signal acquired using flow cytometry (imaged using a FACSCalibur), using concentrations following the manufacturers' instructions. CD117 was detected directly using FITC-conjugated antibody. For detection of FcεRI, LMCs were passively sensitized with IgE overnight, and the bound IgE detected with anti-IgE antibody. Data analysis was done using FlowJo V10 software (BD Biosciences). For detection of FcεRI, LMCs were passively sensitized with IgE overnight, and the bound IgE detected with anti-IgE antibody. Data analysis was done using FlowJo V10 software (BD Biosciences). For degranulation assays, sensitized LMCs were activated through stimulation with 10 ng/mL TNP-BSA for 20 min. Degranulation was determined by measuring secreted and total β-hexosaminidase. The result is reported as a secreted product as a percent of the total β-hexosaminidase. Intracellular calcium mobilization was determined as previously described [25].

ELISA and Western Blotting

IgE-sensitized LMCs were either unstimulated or stimulated with 10 ng/mL TNP-BSA for 30 min, 1, 3, 6, or 24 h, and supernatants collected for ELISA. Antibodies and standards (DuoSet) were purchased from R&D Systems, and ELISAs were conducted according to the manufacturer’s protocol. Control and activated LMCs were lysed in prepared lysis buffer (RIPA) supplemented with HALT™ protease and phosphatase inhibitor cocktail (Thermo Fisher). Protein concentrations were determined in cleared lysates, and 30 μg used for electrophoresis in 10% SDS-polyacrylamide gels. Gels were transferred to polyvinylidene difluoride membranes, blocked with 5% nonfat milk, probed with primary and corresponding secondary antibodies, and detected using an ECL detection system (Western Lightning Plus-ECL; PerkinElmer, Waltham, MA, USA) on a BioMax film (Kodak; Sigma-Aldrich). Blots were scanned and quantified using ImageJ software v1.50.

Real-Time Quantitative Polymerase Chain Reaction

LMCs were either unstimulated or stimulated with 10 ng/mL TNP-BSA, and cell pellets lysed with Trizol (Invitrogen, Thermo Fisher Scientific) for RNA isolation following the manufacturer’s instructions. cDNA synthesis was conducted using 1 μg of RNA and RNA to cDNA EcoDry Premix (Takara Biotech, Mountain View, CA, USA) following the manufacturer’s protocol and thermcycler settings. Samples were prepared for real-time quantitative polymerase chain reaction (RT-qPCR) using the following master mix components: 10 μL molecular grade H₂O (Invitrogen, Thermo Fisher Scientific), 4 μL SsoAdvanced Universal SYBR Green Supermix (Bio-Rad Laboratories), 2 μL 10 μM forward primer, 2 μL 10 μM reverse primer, and 2 μL cDNA. Gene expression was measured using a CFX Connect Real-time System (Bio-Rad Laboratories) and data processed on Bio-Rad CFX Manager 3.1 software. Melt curve analyses were run to determine target specificity. Gene expression levels were normalized to hypoxanthine guanine phosphoribosyltransferase mRNA and analyzed using the ΔΔCt method. Primer sequences are reported in Table 1.

Electrophoretic Mobility Shift Assay

LMCs were unstimulated or stimulated with TNP-BSA, and cell pellets used for nuclear protein isolation. Nuclear protein extracts were obtained using a nuclear extract kit (Active Motif, Carlsbad, CA, USA) according to the manufacturer’s protocol. Double-stranded oligonucleotide probes were labeled using T4 kinase (Life Technologies, Burlington, ON, USA) in the presence of 32P-ATP (PerkinElmer) for 30 min at 37°C. Labeled probes were purified on a Sephadex G-25M column (GE Healthcare, Pittsburgh, PA, USA). Nuclear protein (8 μg) was added to a 10 μL reaction mixture containing binder buffer and supplemented with 1 μg poly-(dI-dC) (GE Healthcare) for 15 min. Labeled and purified probe was added to each reaction mixture and incubated at room temperature for 30 min. Separation by electrophoresis was done on a 6% polyacrylamide gel in 0.5X Tris-boric acid-EDTA buffer. Gels were vacuum dried for 2 h and subjected to autoradiography. The following double-stranded oligonucleotide (Integrated DNA Technologies) was used: NF-kB binding consensus sequence on mouse IL-6 promoter 5′-TTA TCA AAT GTG GGA TTT TCC CAT-3′. Quantification was assessed by densitometric analysis of scanned blots using ImageJ software.
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Reconstitution of MC-Deficient KitW-sh Mice

KitW-sh mice were reconstituted by intradermal injection of LMCs at a density of 25 × 10⁶ cells/mL in 20 μL fresh RPMI 1640 (5 × 10⁵ cells/injection) into the ear pinnae using 31-gauge insulin needles. CnAα+/− LMCs were injected into the right ear pinna, and CnAα−/− LMCs were injected into the left ear pinna. Six weeks later, KitW-sh mice were used in a model of IgE-mediated passive cutaneous anaphylaxis as described below. Nonreconstituted KitW-sh littermates were used as controls. All mice used were sex and age-matched.

IgE-Dependent Passive Cutaneous Reaction

KitW-sh mice reconstituted for 6 weeks with wild-type MCs in right ears, and CnAα+/− MCs in left ears were anesthetized and sensitized with 20 μL of 1 ng/μL anti-DNP IgE mAb (Sigma-Aldrich) using an insulin syringe into the dorsal side of the ear pinnae. Ten microliters of 10 mg/mL DNP-BSA in Evan’s Blue dye (Sigma-Aldrich) was intravenously injected into each sensitized mouse with a 30-gauge needle through the tail vein. The reaction occurred for 30 min before mice were sacrificed, and ear tissues harvested into tubes. Ear tissues were cut into pieces and immersed in 2 mL N, N-dimethyl formamide (Sigma-Aldrich) for 2 h in an 80°C water bath to extract the blue dye. Tubes were centrifuged, and 200 μL of supernant was collected from each to measure absorbance at 620 nm. Only reconstituted mice with successful intradermal injections of LMCs in ear tissues and sensitization were used – if the needle perforates both sides of the ear tissue, there may be possible leakage, and these mice were not used in the analyses. Furthermore, only mice that received successful intravenous tail vein injections on the first attempt were used, to ensure that the same volumes of DNP-BSA in Evan’s Blue dye were administered among the reconstituted mice.

Statistical Analyses

Statistical analyses included 1-way analysis of variance, t tests, and paired t tests as appropriate, and differences were considered significant when p ≤ 0.05. Data displayed in figures are represented as mean ± standard error of the mean.

Table 1. Primer sequences for genes of interest measured using RT-qPCR

| Primer         | Predicted size | Primer sequence 5’−3’          |
|---------------|----------------|--------------------------------|
| TNF forward   | ~174 bp        | CAT CTT CTC AAA ATT CGA GTG ACA A |
| TNF reverse   |                | TGG GAG TAG ACA AGG TAC AAC CC   |
| IL-6 forward  | ~77 bp         | TAG TCC TTC CTA CCC GAA TTT CC   |
| IL-6 reverse  |                | TTT GTC CCT AGC CAC TCC TTC      |
| IL-13 forward | ~108 bp        | CTG TGT CTC TCC CTC TGA CCC      |
| IL-13 reverse |                | GCC AGG TCC ACA CTC CAT ACC       |
| IL-4 forward  | ~77 bp         | CAT GCA CGG AGA TGG ATG TGC      |
| IL-4 reverse  |                | AAG CCC TAC AGA CGA GCT CAC       |
| HPRT forward  | ~229 bp        | CAG ACG ACT AGA ACA CCT GC        |
| HPRT reverse  |                | GCT GGT GAA AAG GAC CTC T         |

RT-qPCR, real-time quantitative polymerase chain reaction; HPRT, hypoxanthine guanine phosphoribosyltransferase; TNF, tumor necrosis factor.

Results

CnAα Wild-Type and Deficient MCs Show Similar Development and Morphology

To determine if CnAα has a role in the development of MCs, CnAα+/− and CnAα−/− LMC cultures were established from neonatal liver cells in media conditioned with mSCF, IL-3, and PGE₂ to promote the maturation and proliferation of liver-derived progenitor cells into MCs. The CnAα gene knockout was confirmed using PCR by the presence of the inserted neomycin cassette used to disrupt a section of the wild-type (WT) gene (Fig. 1a). Characteristics of cultured LMCs, assessed using toluidine blue staining, showed comparable morphology and metachromatic staining for granularity, indicative of similar development in both genotypes of LMCs (Fig. 1b). Furthermore, the kinetics of MC num-

Fig. 1. LMCs from CnAα-deficient mice have similar morphology and development compared to WT in vitro. a Representative genotyping results of WT, CnAα-deficient (KO), and CnAα heterozygous (HZ) samples with labeled DNA ladder. The predicted sizes of CnAα WT and CnAα mutant (neomycin cassette insertion) are 247 bp and 360 bp, respectively. b Representative image using toluidine blue staining of each genotype (original magnification ×100). c, d Flow cytometric detection of surface molecules of WT (left) and CnAα-deficient LMCs (KO, right) sensitized with anti-TNP IgE and then stained with either FITC-conjugated anti-CD117 (c) or anti-IgE antibodies (d). Red peaks are isotype controls. e Intracellular calcium flux was compared between FceRI-mediated activated WT and CnAα-deficient LMCs. BP, base pairs; WT, wild-type; LMC, liver-derived mast cells; CnAα, calcineurin subunit A; FITC, fluorescein isothiocyanate; TNP, trinitrophenol; FceRI, high-affinity receptor for immunoglobulin E; IgE, immunoglobulin E.

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bers and growth between both strains were similar (data not shown). To assess maturation of cultured LMCs, samples were taken for expression analysis of c-kit (CD117) and indirectly, the FcεRI. Analysis by flow cytometry also allows for an objective comparison of the MC morphology based on forward- and side-scatter properties. The forward- and side-scatter properties of MCs of the 2 strains were similar (Fig. 1c, d, left panels), affirming our toluidine blue staining results. Using FITC-conjugated anti-CD117 antibodies, it was deter-

![Graph showing |](image)
determined that \( \text{CnA}^{a-} \) MCs had expression levels of c-kit comparable to \( \text{CnA}^{a+/+} \) MCs (Fig. 1c, right panels). Similarly, using a FITC anti-IgE antibody to indirectly determine expression of FcεRI on IgE-sensitized LMCs, there was comparable expression between \( \text{CnA}^{a+/+} \) and \( \text{CnA}^{a-/-} \) MCs (Fig. 1d, right panels). Furthermore, virtually all gated MCs were positive for both markers in both genotypes of MCs (Fig. 1c, d, middle panels). These results indicate that \( \text{CnA}^{a-/-} \) LMCs are not lacking the receptor for SCF, and additionally, there is no deficiency in the level of FcεRI expression. Understandably, using an anti-IgE antibody is an indirect method of measuring FcεRI, but it also allows us to form a secondary interpretation – that IgE saturation of the receptors between both genotypes is normal and are similar. Antibodies against FcεRI alpha subunits are available but would only give a comparison of surface receptor expression and exclude measurement of IgE binding.

**CnAa Gene Deficiency Does Not Impair Calcium Flux upon FcεRI-Mediated MC Activation**

Calcium signaling precedes and promotes the activation of calcineurin from an inactive to active form and is critical to IgE-dependent signaling cascades. Thus, it is important to assess intracellular calcium fluxes in activated MCs. Sensitized \( \text{CnAa}^{a+/+} \) and \( \text{CnAa}^{a-/-} \) LMCs were incubated with an intracellular calcium indicator, Fura-2 AM, prior to stimulation. The stimulated intracellular calcium flux was found to be unaffected in \( \text{CnAa}^{a-/-} \) LMCs compared to \( \text{CnAa}^{a+/+} \) controls (Fig. 1e). These results indicate that calcium mobilization is not different between WT and \( \text{CnAa}^{a-/-} \) LMCs and signaling leading up to activation of calcineurin is not disrupted.

**CnAa Deficiency Impairs FcεRI-Mediated MC Degranulation in vitro and in Passive Cutaneous Anaphylaxis in vivo**

To assess the effects of \( \text{CnAa} \) deficiency on the early phase of FcεRI-mediated allergic inflammation, sensitized \( \text{CnAa}^{a+/+} \) and \( \text{CnAa}^{a-/-} \) LMCs were stimulated with TNP-BSA for 20 min, and degranulation measured using a β-hexosaminidase assay. \( \text{CnAa}^{a-/-} \) LMCs released approximately 50% less β-hexosaminidase than WT cells (Fig. 2a). Total β-hexosaminidase was determined to assess whether there was a difference that may explain the decreased amount released (Fig. 2b). There was no significant difference between total cell levels of β-hexosaminidase, indicating that release of preformed animal. \( n = 8 \) mice (Experiment was conducted 3 separate times with 3, 3, and 2 mice reaching endpoint, respectively. One mouse did not receive a successful tail vein injection on the first attempt in the final repeat and was omitted). Student’s paired \( t \) test was used for statistical evaluation – \( *p < 0.05 \). WT, wild-type; LMC, liver-derived mast cells; CnAa, calcineurin subunit A; MC, mast cell.
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β-hexosaminidase from granules was impaired in \(CnAα^{−/−}\) LMCs. To determine if the CnAa gene is haplosufficient, LMCs heterozygous for the CnAa gene were also assessed for degranulation, and there was no significant difference when compared to \(CnAα^{+/+}\) (data not shown). Experiments were not continued using the heterozygous genotype. Toluidine blue staining was also conducted on sensitized untreated and sensitized 20-min TNP-stimulated LMCs of both genotypes, to observe granulation before and after MC activation (Fig. 2c). While \(CnAα^{−/−}\) LMCs showed the ability to degranulate, at the light-microscopic level, there were fewer degranulated LMCs and contained more granules than \(CnAα^{+/+}\) LMCs after stimulation, further corroborating our findings that CnAa deficiency resulted in reduced degranulation.

To determine if the in vitro deficiency in degranulation following FcεRI-mediated challenge was observable in vivo, a model of passive cutaneous anaphylaxis was used. Considering that \(CnAα^{−/−}\) mice experience early lethality, in vivo experiments were done on MC-deficient KitW-sh mice reconstituted with WT or \(CnAα^{−/−}\) LMC in ear tissues (no issues with histocompatibility). Importantly, by reconstituting MC-deficient mice, the observations and outcomes upon allergen challenge can be considered as MC-specific. In the model of passive cutaneous anaphylaxis, mice were sensitized with anti-DNP IgE intradermally in the ear for 24 h and then challenged with DNP-BSA in 1% Evan’s blue dye through an intravenous injection. Thirty minutes later, ear tissues were collected from the euthanized mice and Evan’s blue dye extracted as a measure of vascular permeability. Since the entire ear tissue was harvested, histological assessments were not performed on same specimens. There was more blue dye after stimulation with DNP-BSA in the right ears of the mice, for example, reconstituted with WT or \(CnAα^{−/−}\) LMC compared to right ear tissues reconstituted with WT or \(CnAα^{−/−}\) LMCs (Fig. 3a, bottom). Examining multiple animals, there was significantly less vascular permeability in the left ear tissues reconstituted with \(CnAα^{−/−}\) LMCs compared to right ear tissues reconstituted with \(CnAα^{+/+}\) LMCs in the same mouse (Fig. 3b).

These results indicate that there is an impairment in the MC FcεRI-mediated early-phase response to allergen associated with CnAa gene deficiency.

\(CnAα^{−/−}\) LMCs Have Decreased Amounts of Cytokines upon FcεRI-Mediated Activation in vitro

To assess the impact of CnAa deficiency on the late-phase response of FcεRI-mediated allergic inflammation, sensitized \(CnAα^{+/+}\) and \(CnAα^{−/−}\) LMCs were stimulated with TNP-BSA for 1, 3, 6, or 24 h, and cell supernatants collected for detection of cytokines released by MCs including tumor necrosis factor (TNF), IL-4, IL-6, and IL-13. ELISA results for these cytokines showed significantly lower levels of all 4 cytokines in the supernatants at 3, 6, and 24 h of TNP-BSA-stimulated \(CnAα^{−/−}\) LMCs compared to \(CnAα^{+/+}\) (Fig. 4). Furthermore, there seemed to be an overall blunted level of TNF and IL-4 (Fig. 4a, b), yet a steady increase in cytokine levels with time of stimulation in the \(CnAα^{−/−}\) LMCs for IL-6 and IL-13 (Fig. 4c, d). These results indicate that CnAa deficiency leads to an impaired cytokine response, specifically TNF, IL-4, IL-6, and IL-13 in FcεRI-mediated activation in an in vitro system.

To determine if the reduction in TNF, IL-4, IL-6, and IL-13 levels in the supernatants was a result of decreased gene transcript levels or another mechanism, mRNA levels of the 4 cytokines were analyzed using RT-qPCR. The results showed 2 distinct trends – TNF and IL-4 transcript levels were significantly diminished in FcεRI-mediated activated \(CnAα^{−/−}\) LMCs compared to \(CnAα^{+/+}\), whereas there were no significant differences in IL-6 or IL-13 mRNA levels between both genotypes (Fig. 4c, d). These results suggest that CnAa may differentially regulate expression of genes on a cytokine-specific manner in an FcεRI-mediated immune response.

Activation of MAPK Signaling Pathway Family Members Is Not Impaired in \(CnAα^{−/−}\) LMCs

The MAPK signaling cascade family members play critical roles in the transcription of various genes including in inflammatory responses, and activation (phosphorylation) of p38, JNK, and ERK pathways is observed upon FcεRI-mediated stimulation. Western blot results of phosphorylated p38, JNK, and ERK proteins showed comparable activation between \(CnAα^{−/−}\) and \(CnAα^{+/+}\) LMCs (Fig. 5a). Densitometric analyses further confirmed this as there were no significant differences between the 2 genotypes of LMCs (Fig. 5b). These results indicate that MAPK signaling pathway family members...
Fig. 5. CnAα−/− LMCs have similar MAPK family activation but significantly impaired IκBα activation compared to WT. JNK, p38, and ERK phosphorylation were analyzed by Western blotting after FcεRI-mediated activation (a) and quantified using densitometric analyses (b). c CnAα−/− LMCs have significantly less IκBα phosphorylation at 5 and 20 min compared to WT counterparts as quantified by densitometric analysis. A representative blot is shown below the graph. Data are expressed as mean ± SEM (n = 3 separate experiments). Two-way ANOVA was used for statistical evaluation – *p < 0.05. WT, wild-type; LMC, liver-derived mast cells; CnAα, calcineurin subunit A; FcεRI, high-affinity receptor for immunoglobulin E; MAPK, mitogen-activated protein kinase; IκBα, nuclear factor of kappa light polypeptide gene enhancer in B cells inhibitor alpha.
function similarly, regardless of the CnAα gene presence or deficiency. This also raises the notion that these MAPK family members may not play a significant role in regulation of MC degranulation, cytokine synthesis, and release in a CnAα-dependent manner.

Phosphorylation of IκBα Is Impaired in CnAα−/− LMCs in an FcεRI-Mediated Manner

Considering NF-κB signaling is important in the regulation of inflammatory responses and is initiated rapidly upon activation of MCs, the physiological endogenous inhibitor IκBα protein level was measured. Phosphorylation of IκBα leads to its degradation and liberation of NF-κB which translocates to the nucleus and promotes the transcription of various inflammatory genes. Western blot analysis showed significantly less IκBα phosphorylation in CnAα−/− LMCs upon activation in an FcεRI-mediated manner. Specifically, there was significantly impaired phosphorylation of IκBα at 5 and 20 min after TNP-BSA stimulation in CnAα−/− LMCs compared to CnAα+/+ (Fig. 5c). This finding indicates that the NF-κB signaling pathway and associated inflammatory elements are impaired by CnAα deficiency.

Transcription Factor NF-κB Translocation into the Nucleus Is Impaired in FcεRI-Mediated Activated CnAα−/− LMCs

Considering that the inhibitor of NF-κB IκBα had impaired phosphorylation and degradation in CnAα−/− LMCs, the next step was to confirm that nuclear levels of NF-κB were also less in CnAα−/− cells. Banding in the electrophoretic mobility shift assay is the result of protein binding to the radiolabelled oligonucleotides containing a NF-κB binding consensus sequence and thus retarding the probe migration through the gel. When the banding from the blot (Fig. 6a) was quantified using densitometric analysis, there were significantly reduced levels of NF-κB in the nucleus of CnAα−/− LMCs 20 min after stimulation with TNP-BSA compared to CnAα+/+ LMCs (Fig. 6b). This downstream result corresponds with the pattern of phosphorylation and degradation of IκBα seen in Figure 6b, where there was significantly increased activation in CnAα+/+ LMCs from around 5 min to 20 min after MC activation. Furthermore, there is a delayed downward shift in migration of complexes formed between NF-κB proteins and radiolabelled oligonucleotides in the lanes associated with CnAα+/+ LMCs, indicating that there are
increase NF-κB levels in the nucleus in comparison with CnAα−/− LMCs. Collectively, we demonstrated CnAα deficiency resulted in overall decreased degranulation in the allergic response both in vitro and in vivo and decreased secretion of newly synthesized cytokines in vitro, which is likely mediated by impaired NF-κB signaling (Fig. 7).

Discussion

Various types of leukocytes are responsible for the defense of our bodies, requiring these cells to detect and interpret signals from the environment and react in a pre-programmed manner, presumably with an appropriate response. Calcineurin is an important molecule in the flow from interpretation to response. Because of this pivotal role, calcineurin also potentially contributes to undesirable responses leading to harmful consequences. Various calcineurin inhibitors have been used based on anti-inflammatory effects observed in clinical settings, including allergy. Ubiquitous in mammalian tissues and with an evolutionarily conserved catalytic domain, isoforms of calcineurin are likely to have critical functions in cells [26, 27]. Certain isoforms may regulate tissue-specific functions such as development and homeostasis. Our experiments identified a role for the alpha isoform in the context of MC allergy and provide further research directions to expand the limited literature on the contributions of CnAα to disease.

CnAα gene deficiency in mice resulted in changes in maturation in the kidney and epidermal cells but not T- and B-cell growth and development, indicating there are isoform-specific contributions in certain cell types. In particular, others had reported that CnAα−/− mice showed a decrease in vesicle number and protein content of the submandibular glands [23]. Thus, our first assessment was to determine whether CnAα gene deficiency has any effect on structural and granular properties of the MCs. Cultured MCs displayed similar morphology (size and structure, observed microscopically) between both genotypes, and this was confirmed by similar forward-scatter profiles from flow cytometric analysis. Additionally, there was no significant difference found in granules of the MCs, at least measured by β-hexosaminidase content. While MC granules contain multiple factors such as histamine and β-hexosaminidase, both are typically released together from granules [28], and while both are suitable for determining degranulation, the latter is a popular method for assessment [29]. Development and expression of surface receptors c-Kit and FcεRI were also similar between both genotypes. The sum of our findings indicates there is no apparent consequence of CnAα gene deficiency on MC development and maturation, at least among the parameters we measured.
Calcium signaling is integral to FcεRI-mediated signaling and in fact precedes activation of calcineurin, thus it was important to assess intracellular Ca\(^{2+}\) flux in both genotypes. Peak intracellular Ca\(^{2+}\) levels were previously associated with maximal histamine release in a MC line [30], and an absence of sufficient Ca\(^{2+}\) in the cytosol resulted in halting gene transcription as well as rapid shunting of transcription factors back into the cytoplasm to an inactive state [31]. Finding no difference in the pattern and magnitude of the Ca\(^{2+}\) flux, we proceeded to measure functional outcomes following FcεRI activation.

Despite the MCs of the 2 genotypes possessing similar total amounts of β-hexosaminidase, degranulation of preformed mediators was reduced in \(CnAα^{-/-}\) MCs compared to \(CnAα^{+/+}\), as assessed by β-hexosaminidase release in vitro. This difference in degranulation was detectable in the passive cutaneous anaphylactic reaction in mouse ear tissues in the reconstitution experiment. The reduction in the in vivo permeability can be attributed to decreased release of mediators such as histamine from granules as histamine readily promotes vascular permeability [32]. Previously, calcineurin inhibitors were shown to inhibit release of histamine from granules upon activation of MCs in an IgE-dependent manner [33], and now, we build on that finding by showing CnAα is critical to the release of preformed mediators, including by assessing permeability changes driven by these immediate mediators in vivo. Activation of the FcεRI-mediated signaling pathway leads to the activation of a complex of proteins involving Carma1-Bcl10-Malt1 by calcineurin, which may be responsible for downstream signaling through inhibitor of nuclear factor kappa-B kinase 2 (IKK-β) [34, 35]. IKK-β has been identified as a critical signaling component in the pathway leading to degranulation as a deficiency resulted in impaired FcεRI-mediated degranulation of MCs [36]. IKK-β phosphorylates synaptosomal-associated protein 23, which is involved in MC exocytotic mechanisms [36, 37]. Of importance, this mechanism of degranulation is independent of NF-κB activation, which is further corroborated by findings that toll-like receptor signaling (which activates NF-κB) did not induce degranulation in MCs [38]. Although the exact mechanism has not been established, our results demonstrate a significant contribution of CnAα in FcεRI-mediated MC degranulation.

In addition to release of preformed mediators, MCs respond to stimulation through FcεRI with de novo synthesis and secretion of cytokines. Analysis of the secreted mediator levels between stimulated MCs of both genotypes showed cytokine-specific impacts of CnAα. While protein levels of all cytokines examined – TNF, IL-4, IL-6, and IL-13 – were significantly lower in \(CnAα^{-/-}\) than in \(CnAα^{+/+}\) MC supernatants following activation, there were 2 distinct trends in transcript levels of the cytokines. TNF and IL-4 showed parallel low mRNA levels; however, IL-6 and IL-13 gene transcript levels were similar between both genotypes throughout the duration of activation. This may suggest that decreases in IL-6 and IL-13 protein were not due to reduced gene transcription but perhaps other mechanisms dependent on CnAα, such as through interactions with transcription factors or through regulation of secretory pathway mechanisms. For example, in human skeletal muscle cells, calcineurin was found to regulate TNF and IL-6 gene expression differentially, and the addition of calcineurin inhibitors decreased ionomycin-induced IL-6 but not TNF mRNA levels [39]. Possible mechanisms of selective posttranscriptional regulation of these genes also involve mRNA stabilization and translation; however, there is work to be done to identify if there is a role for CnAα in these processes. There may be regulation at the level of mediator release by the soluble NSF attachment protein family as different members have roles in packaging, intracellular trafficking, and extracellular secretion of cytokines and chemokines [40]. A precedent has been published showing calcineurin is crucial in exocytic processes through interactions with soluble NSF attachment family members [41, 42], with the latter implicated in release of mediators by activated MCs.

MAPK family members p38 kinase, JNK, and ERK1/2 are known to contribute to FcεRI-mediated MC activation, and Western blots and respective densitometric analyses showed similarities between the 2 genotypes. This result is comparable to our report using \(Rcan1^{-/-}\) MCs, which exhibited similar phosphorylation of p38, JNK, and ERK1/2 to wild-type levels [16].

The NF-κB signaling pathway is crucial in the transcription of many pro-inflammatory mediators in multiple diseases [43] including allergic inflammation. Our results showed reduced NF-κB in \(CnAα^{-/-}\) MCs in comparison to \(CnAα^{+/+}\). The release and translocation of NF-κB into the nucleus to drive gene transcription are impaired and may be responsible for the observed phenotypes in \(CnAα^{-/-}\) MCs. This outcome again corroborates our study of regulator of calcineurin 1 deficiency, in which loss of the calcineurin inhibitor resulted in significantly greater NF-κB activity and higher gene transcription [16]. Additionally, pharmaceutical calcineurin inhibitors have been shown to reduce NF-κB activity [44]. Activation of the NF-κB signaling axis from calcineurin is possibly through the upstream Carma1-Bcl10-Malt1 complex formation,
which is also important in T-cell receptor activation [45]. This was corroborated by other research finding a role for calcineurin in transient dephosphorylation of Bcl-10 to be necessary for NF-κB activation in T H 1 cells [34].

Considering none of our measures were absolutely abolished, there was residual calcineurin activity, likely due to the β isoform, so determining the role of this isoform remains to be done. The β isoform has been regarded as the predominant isoform in lymphocytes and plays a critical role in immune settings [10]. Meanwhile, it is evident that CnAα definitively and CnAβ likely contribute to the FcεRI-mediated immune response.

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Statement of Ethics

Protocols (18-003 and 18-004) were approved by the University Committee on Laboratory Animals, Dalhousie University, Canada, in accordance with the guidelines of the Canadian Council of Animal Care.

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Conflict of Interest Statement

The authors declare no conflicts of interest.

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Author Contributions

E.L. contributed to conceptualization, generation, and formal analysis of experimental data for this manuscript. Z.P. provided guidance and assistance with experimental assays. A.W.S. and T.-J.L. provided concept design, guidance, and supervision for this research project. E.L. and A.W.S. prepared and edited the manuscript. Address all correspondences to Dr. Andrew W. Stadnyk at astadnyk@dal.ca.

Data Availability Statement

All data generated or analyzed during this study are included in this article. Further inquiries can be directed to the corresponding author.
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