Transplacental Innate Immune Training via Maternal Microbial Exposure: Role of XBP1-ERN1 Axis in Dendritic Cell Precursor Programming

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We recently reported that offspring of mice treated during pregnancy with the microbial-derived immunomodulator OM-85 manifest striking resistance to allergic airways inflammation, and localized the potential treatment target to fetal conventional dendritic cell (cDC) progenitors. Here, we profile maternal OM-85 treatment-associated transcriptomic signatures in fetal bone marrow, and identify a series of immunometabolic pathways which provide essential metabolites for accelerated myelopoiesis. Additionally, the cDC progenitor compartment displayed treatment-associated activation of the XBP1-ERN1 signalling axis which has been shown to be crucial for tissue survival of cDC, particularly within the lungs. Our forerunner studies indicate uniquely rapid turnover of airway mucosal cDCs at baseline, with further large-scale upregulation of population dynamics during aeroallergen and/or pathogen challenge. We suggest that enhanced capacity for XBP1-ERN1-dependent cDC survival within the airway mucosal tissue microenvironment may be a crucial element of OM-85-mediated transplacental innate immune training which results in postnatal resistance to airway inflammatory disease.

Keywords: innate immune training, myelopoiesis, immunomodulator, XBP1, dendritic cell, transplacental, myeloid progenitor, OM-85

INTRODUCTION

The neonatal period represents a time of high risk for infection-related morbidity/mortality resulting from the combined effects of maturational deficiencies in both anti-microbial defense mechanisms that mediate pathogen recognition and elimination, and in the accompanying regulatory mechanisms required for calibration of these responses to minimize inflammatory
damage to host tissues (1, 2). With respect specifically to the lung tissue microenvironment, a crucial factor governing the kinetics of postnatal acquisition of immune competence is the rate of development of the airway mucosal cDC network which controls local immune surveillance (3, 4).

In addition to increased susceptibility to infectious diseases, the seeds for development of a range of non-communicable diseases exemplified by asthma and aero-allergies are also frequently sewn during this early postnatal window period (5), suggesting that maturational deficiencies in immune function(s) may also be risk factors in this context. In this regard, maternal immune perturbations have been acknowledged to significantly influence fetal immune development, but the underlying mechanisms remain poorly characterized (6). Epidemiological data from studies on traditional farming families in Europe and USA suggesting that benign environmental microbial exposures of mothers during pregnancy can promote prenatal immune maturation within their offspring, leading to reduced susceptibility to postnatal development of respiratory inflammatory diseases (7, 8), have stimulated wide-spread interest in this issue. This capacity for microbial exposures to modulate immune development is consistent with the paradigm of “immune training”, whereby exposure to certain classes of microbial stimuli can alter the long-term functional state of innate immune cells, occurring at the progenitor level in the bone marrow (BM) (9–11), leading to optimized peripheral immune responsiveness to other unrelated microorganisms (12). With this in mind, there is growing interest in the concept that immune training can be therapeutically harnessed (13), particularly during prenatal development, to enhance immunocompetence within the offspring (14).

We recently reported that oral treatment of pregnant mice with the microbial-derived immunomodulator OM-85 reduces susceptibility of their offspring to the development of Th2-driven allergic airways inflammation, and identified myeloid progenitors in the offspring BM (which supply precursor DC to eventually populate mucosal DC networks) as a major target for maternal treatment effects (15). In the study presented here, we employed transcriptomic profiling to characterize gene networks activated in fetal BM (BM) as a result of maternal OM-85 treatment, and identify the principal treatment targets as immunometabolic pathways supplying cellular cholesterol essential for rapid expansion of myeloid precursor compartments, and which have previously been recognized as hallmarks of classical immune training-associated gene signatures. We additionally identify activation of the XB1P-ERN1 signalling axis in the cDC precursor compartment, which has previously been associated with survival-under-stress, especially within the lung mucosal microenvironment.

**METHODS**

**Animals**

Specific pathogen-free BALB/c mice were purchased from the Animal Resource Centre (Murdoch, Western Australia, Australia). All mice were housed under specific pathogen-free conditions at the Telethon Kids Institute Bioresources Centre.

**Time-Mated Pregnancies**

Female BALB/c mice 8–12 weeks of age were time-mated with male BALB/c studs 8–26 weeks of age. Male studs were housed individually with 1–2 females overnight. The detection of a vaginal plug the following morning was designated gestation day (GD) 0.5.

**Maternal OM-85 Treatment**

OM-85 (OM Pharma) is an endotoxin-low lyophilized extract containing a cocktail of TLR ligands derived from 8 major respiratory tract bacterial pathogens (*Haemophilus influenzae*, *Streptococcus pneumoniae*, *Streptococcus pyogenes*, *Streptococcus viridians*, *Klebsiella pneumoniae*, *Klebsiella ozaenae*, *Staphylococcus aureus* and *Neisseria catarrhalis*) (16, 17). Based on previously optimized dosing concentrations (18, 19), time-mated pregnant BALB/c mice received daily oral feeding of lyophilized OM-85 reconstituted in phosphate-buffered saline (PBS; prepared in-house) via pipette at a concentration of 400mg/kg body weight for the second half of gestation (GD9.5 – 17.5). Control pregnant mice were left untreated for the duration of the study, as based on our previously published studies which demonstrated no difference between PBS (vehicle)-treated and untreated mothers (18). All pregnant mice were handled equivalently throughout the duration of the study. All maternal treatment was performed with a single batch of OM-85 (batch# 1812162).

**Tissue Collection**

Pregnant BALB/c mice were sacrificed 24 h after the final OM-85 dose at GD18.5. Both horns of the uterus were removed and fetuses sacrificed by decapitation. Fetal hind legs (cleaned of excess tissue) were removed and long bones (femur and tibia) collected. Fetal long bone samples for flow cytometry were collected into cold PBS + 0.1% bovine serum albumin (BSA) and stored on ice. Fetal long bone samples for transcriptomic analysis were collected into RNAlater® stabilization solution (Sigma-Aldrich). Samples collected into RNAlater® were stored overnight at 4°C, then transferred to 1.5ml Eppendorf tubes (Eppendorf) and frozen at −80°C for future transcriptome profiling. All fetal samples were kept as individuals and not pooled. Dead fetuses were excluded from the study.

**Single-Cell Suspension Preparation**

Fetal long bones were prepared by mincing with a scalpel followed by enzymatic digestion, as previously detailed (15). Briefly, minced bones were resuspended in 10ml GKN (11mM D-glucose, 5.5mM KCl, 137mM NaCl, 25mM Na2HPO4; prepared in-house) + 10% fetal calf serum (FCS; Serana) with collagenase IV (Worthington Biochemical Corp.) and DNase (Sigma-Aldrich) at 37°C under gentle agitation for 60 min. Digested whole bone homogenates were filtered through sterile cotton wool columns (5ml syringe containing cotton wool; prepared in-house) coated with FCS to remove debris, centrifuged and resuspended in cold PBS for total cell counts.

**Flow Cytometry**

Fetal whole bone single-cell suspensions (prepared above) were used for all immunostaining. Panels of monoclonal antibodies
(purchased from BD Biosciences unless otherwise stated) were developed to enable phenotypic characterization of committed myeloid cells: CD3-FITC (clone 17A2), CD11b-BV510 (clone M1/70), CD11c-BV711 (clone HL3), CD19-APC-H7 (clone 1D3), Gr-1-Biotin (clone RB6-8C5), CD45R/B220-PerCP-Cy5.5 (clone RA3-6B2) NKp46-PE-Cy7 (clone 29A1.4; BioLegend), SIRPα-APC (clone P84; BioLegend), I-A/I-E-BV421 (clone M5/114.15.2; BioLegend), F4/80-BV785 (clone BM8; BioLegend), Viability-AF700, Streptavidin-BV605; hematopoietic stem and progenitor cells: CD2-Biotin (clone RM2-5), CD3-Biotin (clone 145-2C11), CD4-Biotin (clone GK1.5), CD5-Biotin (clone 53-7.3), CD8α-Biotin (clone 53-6.7), CD19-Biotin (clone 1D3), CD45R/B220-Biotin (RA3-6B2), Gr-1-Biotin (clone RB6-8C5), Ter119-Biotin (clone TER-119), CD16/32-PerCP-Cy5.5 (clone 2.4G2), CD34-FITC (clone RAM34), IL-7R-APC (clone SA011F11; BioLegend), NKG2D-PE-Cy7 (clone SB/199), Flt3-PE (clone A2F10.1), c-Kit-APC-Cy7 (clone 2B8), Sca-1-BV510 (clone D7), CX3CR1-APC (clone SA011F11; BioLegend), NKG2D-BV711 (clone CX5), Viability-AF700, Streptavidin-BV605 and XBP1s-expressing bone marrow cells: CD3-FITC (clone 17A2), CD11b-BV510 (clone M1/70), CD11c-AF700 (clone HL3), CD19-APC-H7 (clone 1D3), I-A/I-E-AF647 (clone M5/114.15.2), CD45R/B220-PE-CF594 (RA3-6B2), Gr-1-Biotin (clone RB6-8C5), NKp46-PE-Cy7 (clone 29A1.4; BioLegend), F4/80-BV785 (clone BM8; BioLegend), XBP1s-BV421 (clone Q3-695), Streptavidin-BV605. Intracellular staining for XBP1s was performed using an intracellular Foxp3/Transcription factor staining buffer kit (eBioscience). Data acquisition was performed on a 4-laser LSRFortessa (BD Bioscience). All samples were kept as individuals and not pooled. Immune cell phenotypic characterization was performed using FlowJo software (version 10.1, Tree Star). Fluorescence minus one (FMO) staining controls were used for all panels where necessary (Supplementary Figure 1). Flow cytometry data quality was based on primary time gates to ensure appropriate laser delay (pre-determined by automated CS&T) during sample acquisition.

Flow Cytometric Statistical Analyses
Statistical analysis and graphing was performed using GraphPad Prism (GraphPad software; version 7.0a). Statistical significance of p<0.05 was considered significant. Unpaired, two-tailed Student’s t-test or Mann Whitney U test were used based on distribution of the data as determined by D’Agostino-Pearson omnibus normality test. Internal correlation within the untreated and OM-85 treated groups was assessed by controlling for family clustering using Generalized Estimation Equation (20) or Wilcoxon Rank-Based Test for Clustered Data (arXiv:1706.03409) as based on distribution of the data. Significance of the findings were not influenced by family clustering. Part of the cDC flow cytometry data presented in Figure 1 and MDP data presented in Figure 2 has been published in a forerunner manuscript (15).

Transcriptome Profiling (RNA-Seq)
Tissue Preparation, RNA Extraction, and Transcriptome Profiling
Fetal bone marrow samples were homogenized using a rotor-star homogenizer (Qiagen) and total RNA extracted via TRIzol (Invitrogen), followed by clean-up using RNeasy MinElute Cleanup Kit (Qiagen). RNA integrity was determined using an Agilent 2100 Bioanalyzer (Agilent Technologies; RIN: 10 ± 0 (mean ± SD)). One microgram total RNA (n = 32) was shipped on dry ice to the Australia Genome Research Facility (AGRF) for library preparation (TruSeq Stranded mRNA Library Prep Kit, Illumina) and sequencing (Illumina HiSeq2500, 50-bp single-end reads, v4 chemistry).

RNA-Seq Data Analysis
Pre-Processing and Exploratory Data Analysis
RNA-seq data was analysed in the R environment for statistical computing. Sequencing data quality control (QC) was performed with the Bioconductor package Rqc (21). Sequencing reads were aligned to the reference murine genome (mm10) using Subread and summarized at the gene level using featureCounts (22). Genes with <500 total counts across the data were removed from the analysis. Sample QC was performed by analysing the distribution of the raw read counts to check for sample outliers using boxplots, relative log-transformed expression (RLE) plots and principal component analysis (PCA), before and after global-scale median normalization. Differential expression analysis: Differentially expressed genes (DEG) were identified employing the DESeq2 package (23). DESeq2 utilizes a negative binomial distribution model, with a False Discovery Rate adjusted P-value for multiple comparisons. Genes were deemed significant with an adjusted P-value < 0.1. Pathways analysis: Pathways enrichment analysis was performed using the InnateDB database (24) with Benjamini & Hochberg adjusted P-value ≤ 0.05 deemed significant. Upstream regulator analysis: Ingenuity Systems Upstream Regulator Analysis (25) was employed to identify putative molecular drives of the DEG patterns. Significance was determined by activation Z-score ≥ 2 and P-value of overlap ≤ 0.05.

Study Approval
All animal experiments were formally approved by the Telethon Kids Institute Animal Ethics Committee, operating under the guidelines developed by the National Health and Medical Research Council of Australia for the care and use of animals in scientific research.

RESULTS
Maternal OM-85 Treatment Selectively Accelerates Functional Maturation of cDCs in Fetal Bone Marrow
To elucidate the mechanisms-of-action of maternal OM-85 treatment, we examined the in utero fetal response at gestation day (GD) 18.5 (Figure 1A), 2 days prior to expected natural term delivery. Primary observations identified a significant increase in the cellularity of fBM following maternal OM-85 treatment as compared to fBM from untreated mothers (data not shown). Targeted phenotypic analysis of the fBM myeloid compartment
using multicolor flow cytometry (Supplementary Figure 2) revealed significant expansion of the total dendritic cell (DC) pool in fetuses from OM-85 treated mothers as compared to equivalent fetal samples from untreated mothers (Figure 1B). Further characterization of the BM DC response demonstrated that this increase was restricted to the CD11b+B220-CD11c+Gr-1-SIRPa+I-A/I-E+ conventional DC (cDC) subset as previously described (15), with no parallel changes observed in CD11b-B220+CD11c+Gr-1+I-A/I-E+ plasmacytoid DC (pDC) (Figure 1B). These cDC-specific changes in fBM mirror our recent observations of increased cDC yields from BM cultures and peripheral lung from the offspring of OM-85-treated mothers in the early postnatal period (15). We next turned our attention to fetal DC maturation state as determined by surface I-A/I-E expression. As shown in Figure 1C, maternal OM-85 treatment for the last half of gestation enhanced I-A/I-E expression on cDC in fBM when compared to fBM cDC from untreated mothers. Collectively, these observations suggest that transplacental signals generated at the feto-maternal interface following OM-85 treatment during pregnancy can "train" the developing fetal immune system via promoting the development of a fBM cDC compartment exhibiting a phenotype associated with enhanced functional competence.

**Expansion of Fetal Bone Marrow Myeloid Progenitor Subsets Following Maternal OM-85 Treatment**

Previous studies from our laboratory have additionally identified postnatal expansion of BM myeloid progenitor (MP) cell populations as an effect of maternal treatment with OM-85 during pregnancy (15). These findings mirror that of recent studies which identified modulation of BM MP as an important component of conventional immune training mediated by both β-glucan (10, 26) and Bacillus Calmette-Guérin (BCG) (11). Based on these findings, we hypothesized that the enhanced cDC population within fBM following maternal OM-85 treatment would also be accompanied by concomitant upregulation of upstream MP subsets. Although the linear commitment model of myeloid progenitor subsets remains controversial (27–29), using this approach for our hierarchical flow cytometric analysis of fBM (Figure 2A; Supplementary Figure 3) demonstrated a significant increase in total Lin IL-7Ra c-Kit'Sca-1' MP (Figure 2B), Lin IL-7Ra c-Kit'Sca-1'CD16/323CD34+ granulocyte-macrophage progenitor (GMP; Figure 2C) (30, 31) and Lin IL-7Ra c-Kit'Sca-1' CD16/323CD34+CX3CR1 Flt-3' macrophage-dendritic cell progenitor (MDP; Figure 2D) (32, 33) populations.
within the BM compartment following maternal OM-85 treatment, when compared to fBM from untreated mothers. However, no changes were observed in the Lin-IL-7Rα-c-Kit+Sca-1+Flt-3+ common lymphoid progenitor (CLP) (34, 35) or Lin-IL-7Rα-c-Kit+Sca-1+NKG2D+ pre-natural killer cell progenitor (pre-NKp) (36, 37) populations following maternal OM-85 treatment (Supplementary Figure 4). Consistent with the findings in Figures 1B, C, these data provide further evidence that maternal OM-85 treatment selectively modulates the offspring BM myeloid lineage in utero, beginning at the early-stage myeloid progenitor level through to the terminal cDC populations which are responsible for seeding peripheral tissues during early postnatal life to provide local DC-mediated immune surveillance.

Maternal OM-85 Treatment Activates Key Regulators of the UPR Pathway in Fetal Bone Marrow

To gain further insight into the molecular mechanisms underpinning the maternal OM-85-treatment effects, we employed transcriptomic profiling of fBM cells. Comparison of the transcriptomic profiles in the treated versus untreated groups
indicated that maternal OM-85 treatment resulted in 152 differentially expressed genes (DEG) in fBM (119 upregulated, 33 downregulated; Figure 3A, Supplementary Table 1). We then interrogated the DEG for enrichment of biological pathways employing InnateDB (24), focusing on the upregulated DEG response given the limited number of downregulated DEG identified. In fBM, upregulated DEG were enriched for genes involved in multiple aspects of protein metabolism, the endoplasmic reticulum (ER) stress response, the unfolded protein response (UPR), cholesterol biosynthesis and lipid metabolism (Figure 3B; Supplementary Table 2).

Upstream regulator analysis was then performed to identify putative molecular drivers of all observed DEG. The data revealed X-box binding protein 1 (XBP1), a transcription factor central to the UPR (38) and crucial in the development, survival and function of multiple cell types including plasma cells (39), eosinophils (40), natural killer (NK) cells (41), T-cell subsets (42) and DCs (43, 44), as the most strongly activated molecular driver within fBM associated with maternal OM-85 treatment effects ($P$-value = 3.81x10^{-17}, Z-score = 4.427, Figure 3C; Supplementary Table 3), and consistent with this, its downstream target, the canonical UPR sensor Activating Transcription Factor 6 beta (ATF6b) (45, 46) was upregulated (Supplementary Table 1 and Supplementary Table 3). Additionally, Endoplasmic Reticulum To Nucleus Signalling 1 (ERN1) was identified as an activated driver gene within the fBM following maternal OM-85 treatment ($P$-value = 3.32x10^{-6}, Z-score = 2.156; Figure 3C; Supplementary Table 3). Identification of ERN1 is crucial given that during the ER stress response, this gene encodes the ER stress sensor protein inositol-requiring enzyme 1 (IRE1ε), responsible for the unconventional cleavage of a 26 nucleotide fragment from Xbp1 mRNA, resulting in the generation of the active spliced variant of XBP1 (XBP1s) and enabling it to function as a potent molecular driver.
transcription factor within the UPR signalling pathway (46), as evidenced by XBP1 being identified as a downstream target gene of activated ERN1 (Supplementary Table 3). Collectively, these findings suggest that activation of the UPR pathway may be a central component of the immune training mechanism induced in fBM as a result of maternal OM-85 treatment. In addition to UPR pathway drivers, maternal OM-85 treatment also resulted in the upstream activation of multiple drivers central to immunometabolic pathways involved in cellular cholesterol homeostasis, including Sterol Regulatory Element Binding Transcription Factor 1 (SREBF1; P-value = 1.95x10⁻⁶, Z-score = 3.056) and 2 (SREBF2; P-value = 2.95x10⁻⁹, Z-score = 2.745) and Sterol Regulatory Element Binding Protein Cleavage-Activating Protein (SCAP; P-value = 1.52x10⁻¹⁰, Z-score = 2.949; Figure 3C; Supplementary Table 3). This was associated with upregulation of their downstream target Low-Density Lipoprotein Receptor (LDLR; Supplementary Table 1 and Supplementary Table 3), while Insulin Inducible Gene 1 (INSIG1) was inhibited in fBM following maternal OM-85 treatment (P-value = 5.81x10⁻⁸, Z-score = −2.931; Figure 3C; Supplementary Table 3).

Additional candidate drivers in fBM following maternal OM-85 treatment included CD38, IL5, and an array of microRNAs (miR) (Figure 3C; Supplementary Table 3) recognized principally in the context of cancer-associated functions (47–49). Of note, miR-149-3p (strongly downregulated in Figure 3C) has been shown to negatively regulate Toll-like receptor (TLR) 4 expression in murine monocytic cells in vitro (50) and it is possible that other miRs may have similar (but as yet undefined) innate immune regulatory functions (51). The relevance of finding TLR4 upregulation in BM-derived myeloid cells in this model merits further investigation. Likewise, the identification of the T-cell activation-associated markers CD38 and IL-5 suggests possible contributions from activated T-cells, and these possibilities will also be addressed in follow up studies.

DISCUSSION

In this study, we have characterized the response of the fBM myeloid cell compartment to maternal treatment with the microbial-derived innate immune modulator OM-85 during pregnancy. We demonstrate that, consistent with our previous reports in 6-week-old offspring (15), maternal OM-85 treatment expanded the baseline pool of GMP and MDP in fBM. Furthermore, we show for the first time the maternal OM-85-induced modulation of metabolic pathways within fBM responsible for cellular cholesterol homeostasis, with specific activation of SREBF1, SREBF2, SCAP and LDLR. In this regard, SREBF1 and SREBF2 are responsible for encoding sterol regulatory element binding proteins (SREBP) which play a central role in cellular metabolism by controlling the synthesis of cholesterol and other membrane lipids in the Golgi of mammalian cells, with SREBP2 (encoded by SREBF2) regarded as the “master regulator” of cellular cholesterol biosynthesis (53).
Furthermore, SCAP is central to this process by acting as a protein chaperone, mediating the transport of SREBP from the ER to the Golgi where it can promote transcription of 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase, the rate-limiting enzyme of the mevalonate pathway responsible for intracellular cholesterol biosynthesis (53). In parallel with this metabolic pathway, LDLR regulates cellular uptake of low-density lipoprotein (LDL), the upregulation of which increases downstream cholesterol accumulation within the cell and is required for the proliferation of hematopoietic stem and progenitor cells (54). It is now recognized that myelopoiesis within the BM is heavily reliant upon an increased demand for cellular cholesterol and enhanced cholesterol biosynthesis (55–57), and we therefore postulate that upregulation of this immunometabolic pathway following maternal OM-85 treatment is in-part responsible for the expansion of MP, GMP and MDP observed within fBM. Further reinforcing the importance of enhanced cholesterol biosynthesis, INSIG1, an ER membrane protein that prevents trafficking of SCAP/SREBP complexes to the Golgi and thereby terminates cholesterol biosynthesis by restricting HMG-CoA reductase-mediated activation of the mevalonate pathway (53, 58), was inhibited in fBM following maternal OM-85 treatment. Together, these findings parallel previous studies demonstrating that activation of cellular cholesterol biosynthesis and resultant expansion of BM MP is a hallmark of classical β-glucan-mediated central immune training, and suggests a common role for the mevalonate pathway in these immune training mechanisms (10, 59). We further demonstrate that transplacental mechanisms promoting maternal OM-85-induced immune training within fBM involve a dynamic process comprising upregulation of immunometabolic pathways that provide key rate-limiting metabolites required for myelopoiesis and subsequent expansion of MP, GMP and MDP, and associated inhibition of negative feedback loops responsible for arresting cholesterol biosynthesis.

Downstream of the fBM progenitor response, maternal OM-85 treatment selectively amplified the overall abundance of fBM cDC, along with enhancing the concomitant functional maturation of these cDC as demonstrated by upregulated I-A/I-E (MHC Class II) expression. This BM population is the source of the precursors which subsequently seed the airway mucosal DC network that progressively develops between birth and weaning (3, 4). It is noteworthy that the cDC which initially seed this network postnatally are MHC-IIlow (reflecting their functionally immature status) relative to the high-level expression seen at later ages (3), and the findings above in fetal cDC from the treated group may collectively explain the accelerated postnatal establishment and the enhanced functional maturation of this network observed in their offspring (15). This DC network plays an essential “gatekeeper” role in immune surveillance of airway surfaces, and hence in protection against both allergic and infectious diseases in the respiratory tract (60, 61), and its relative paucity and reduced functionality during infancy may be an important contributor to increased susceptibility to these diseases during this life phase.

Earlier studies from our group also identified unique features of the population dynamics of this lung cDC network which distinguishes it from comparable populations in other tissues, notably the exceptionally rapid baseline turnover rate of individual cells within the network, ~85% of which are replaced every 24–36 h (62), with emigration to draining lymph nodes (bearing samples of locally acquired antigens) balanced by recruitment of replacements from BM. Moreover, once development of functional competence is complete (4), this network develops capability for rapid expansion to up to 5-fold baseline density in the face of acute challenge with airborne pro-inflammatory irritant, allergic or microbial stimuli (63), the latter response exhibiting kinetics that rival neutrophils (64, 65). These unique population dynamics suggest that even at baseline, mechanisms that promote lung cDC survival are likely to play a crucial role in the capacity of the network to perform its immune surveillance functions which require onward migration to downstream lymph nodes and subsequent interaction with T-cells as opposed to antigen presentation in situ (60); moreover during prolonged/severe events exemplified by severe viral infections, the added effects of cDC injury resulting from inflammation-associated ER stress (66) would place further pressure on survival times.

In this regard, our transcriptomic analyses of fBM from offspring of OM-85-treated mothers also identified upregulated expression of XBP1, ATF6β and ERN1 (encoding IRE1α), key components of the XBP1-ERN1 signalling axis and critical regulators of the UPR pathway (67) which mitigates the effects of ER stress, and moreover we localized upregulated production of active XBP1s protein to cDC precursors. While this enhanced XBP1s cDC precursor pool constitutes a small population within the totality of the fBM, individual precursor cells have a remarkable ability to generate vast pools of mature DC (68), and as such heavily influence the downstream function of peripheral tissue DC populations. Taken together with recent findings on the role for IRE1α-XBP1 signalling and the downstream transcription factor XBP1s in DC development and function (43, 44), these results suggest a central role for the XBP1-ERN signalling axis in this OM-85-mediated immune training process.

In further support of this suggestion, other studies demonstrate a significant reduction in CD11c+ cells (mirroring that of our pre-cDC phenotype) within XBP1−/− BM cultures, whilst forced overexpression of XBP1s in XBP1−/− DC precursors conversely rescues and subsequently drives expansion of the DC pool in vitro (43). It is also pertinent to note that others have reported that the cDC population in the lung mucosa is differentially reliant upon XBP1 expression for survival at baseline relative to cDC from other tissue sites (66), which may be a direct reflection of the uniquely high turnover rates of cDC within the airway mucosal microenvironment (62).

Collectively, the data presented here indicate that OM-85 likely operates as an immune training agent, employing cellular and immunometabolic mechanisms previously reported in independent model systems (10, 11, 69), with the additional capacity to act transplacentally via the fBM. Furthermore, we go beyond the currently known features of innate immune training...
to describe involvement of the XBP1-ERN1 signalling axis. Moreover, classical β-glucan- and BCG-mediated immune training has traditionally focused on prototypic innate effector cell populations (monocytes/macrophages/natural killer cells) resulting in enhanced resistance to bystander pathogens via the upregulation of pro-inflammatory responses exemplified by tumor necrosis factor-α (TNF-α), interferon-γ (IFN-γ), IL-6 and IL-1β production (70–72) and emergency granulopoiesis-mediated neutrophil influx (69). However, studies are now beginning to recognize that immune training can also occur in DC populations, resulting inter alia in epigenetic reprogramming of pro-inflammatory cytokine responses (73). We have recently extended these observations to include OM-85-mediated training effects on key immunoregulatory functions in both pregnant mice and their offspring, including effects on both cDC and pDC populations and downstream T-regulatory cells, which are collectively associated with enhanced resistance of both mothers and offspring to the pro-inflammatory effects of bacterial, viral and allergic stimulation (15, 18). Of note, similar transplacental immune training-like effects targeting immunoregulatory mechanisms in offspring have been reported in relation to pregnant maternal exposure to extracts from Acinetobacter lwofii (74) and Helicobacter pylori (75), suggesting that the phenomenon reported here may be generalizable, and may point towards a novel approach to mitigation of disease risk in the age group that is in greatest need of protection.

DATASET AVAILABILITY STATEMENT

The dataset generated for this study can be found in the Gene Expression Omnibus repository: https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE140143.

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ETHICS STATEMENT

The animal study was reviewed and approved by Telethon Kids Institute Animal Ethics Committee.

AUTHOR CONTRIBUTIONS

KM, PH, and DS designed the study. KM, MB, NS, and J-FL-J performed the experiments. KM, AJ, MB, and DS analyzed the data. PS and AB contributed to the project design and discussions on data interpretation. KM, PS, PH, and DS wrote the manuscript. 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/fimmu.2020.601494/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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