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Xiaofang Tang, …, Xinhua Lin, Jeffrey A. Whitsett


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EMC3 coordinates surfactant protein and lipid homeostasis required for respiration

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Adaptation to respiration at birth depends upon the synthesis of pulmonary surfactant, a lipid-protein complex that reduces surface tension at the air-liquid interface in the alveoli and prevents lung collapse during the ventilatory cycle. Herein, we demonstrated that the gene encoding a subunit of the endoplasmic reticulum membrane complex, EMC3, also known as TMEM111 (Emc3/Tmem111), was required for murine pulmonary surfactant synthesis and lung function at birth. Conditional deletion of Emc3 in murine embryonic lung epithelial cells disrupted the synthesis and packaging of surfactant lipids and proteins, impaired the formation of lamellar bodies, and induced the unfolded protein response in alveolar type 2 (AT2) cells. EMC3 was essential for the processing and routing of surfactant proteins, SP-B and SP-C, and the biogenesis of the phospholipid transport protein ABCA3. Transcriptomic, lipidomic, and proteomic analyses demonstrated that EMC3 coordinates the assembly of lipids and proteins in AT2 cells that is necessary for surfactant synthesis and function at birth.

Introduction

Pulmonary surfactant is a complex mixture of lipids and associated proteins synthesized by alveolar type 2 (AT2) cells in the peripheral lung. Surfactant is required at birth and throughout postnatal life to reduce surface tension at the air-liquid interface in the alveoli. Lack of surfactant causes atelectasis and respiratory distress syndromes in premature infants (RDS) and adults (ARDS) (1). Surfactant proteins, and a phospholipid transport protein, ABCA3, are required for the packaging of phospholipids into specialized organelles termed lamellar bodies (LBs) by AT2 cells and for the function of surfactant in the alveoli. Mutations in the ABCA3, SFTPB, and SFTPC genes cause severe lung disease in newborn infants and children, usually presenting with respiratory failure soon after birth (2–4). The folding, proteolytic processing, and intracellular trafficking of surfactant proteins in LBs are critical for the production of pulmonary surfactant. In this report, we tested whether EMC3 plays a role in the assembly of surfactant lipids and proteins in AT2 cells in the neonatal mouse lung. We show that, in spite of normal lung morphogenesis, deletion of Emc3 in embryonic lung epithelial cells led to respiratory distress and neonatal lethality related to surfactant deficiency. Furthermore, we demonstrate that EMC3 controls the processing and routing of surfactant proteins SP-B and SP-C as well as the biogenesis of the phospholipid transporter protein ABCA3. Therefore, we have identified EMC3 as a critical component of an ER complex that mediates the processing and trafficking of surfactant
associated proteins and lipids by AT2 cells and is required for lung function and respiration at birth.

Results

Conditional deletion of Emc3 in respiratory epithelium causes surfactant deficiency and neonatal respiratory failure. To assess the role(s) of EMC3 in lung epithelial cells, an Emc3fl/fl allele was produced and mated to Shh-Cre–transgenic mice to delete Emc3 in respiratory epithelial cells prior to birth (Figure 1, A and B). While Shh-Cre;Emc3fl/+ mice survived normally, all Shh-Cre;Emc3fl/fl mice died of respiratory failure immediately after birth (Table 1). Cre-mediated excision in Shh-Cre;Emc3fl/fl (referred to as Emc3-cKO hereafter) mice reduced Emc3 mRNA levels in whole-lung homogenates and isolated EpCAM+ (CD326+) epithelial cells at E18.5, the day before birth (Figure 1C and Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI94152DS1). Western blot analysis demonstrated decreased EMC3 protein levels (Figure 1D). Deletion of Emc3 decreased the other EMC proteins we examined including EMC1, 4, 7, and 10 but did not influence their RNAs, indicating the essential role of EMC3 in stabilizing the entire EMC complex (Figure 1D, and Supplemental Figure 1, A and C). Without any observable malformations, Emc3-deficient pups remained cyanotic and failed to inflate their lungs in spite of vigorous respiratory efforts, while control (Emc3+/+) littermates were well oxygenated and survived (Figure 2, A–D). Histological analysis of lung sections at E18.5 showed no abnormalities in lung morphogenesis (Figure 2, E and F). Differentiation of AT1 cells, indicated by podoplanin (PDPN), AGER, and HOPX staining, was undisturbed (Supplemental Figure 2), supporting the concept that defective surfactant homeostasis rather than a developmental delay may underlie the respiratory failure seen in the Emc3-cKO mice. Ultrastructure of AT2 cells in Emc3-mutant lungs lacked normal LBs and instead contained small, poorly lamellated vesicles, while the ultrastructure of other intracellular organelles was unchanged (Figure 2, G and H, and Table 2). Given the essential function of LBs in surfactant packaging, secretion, and function, the abnormal LB morphology associated with Emc3 deletion demonstrated that EMC3 played an important role in surfactant biosynthesis before birth.

Table 1. Postnatal death in Emc3-cKO mice

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Progeny at weaning (n = 80)</th>
<th>Progeny at E18.5 (n = 149)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Emc3+/+</td>
<td>Emc3fl/fl</td>
</tr>
<tr>
<td>Expected</td>
<td>25%</td>
<td>25%</td>
</tr>
<tr>
<td>Shh-Cre;Emc3fl/+</td>
<td>32.5% (26)</td>
<td>35.0% (28)</td>
</tr>
<tr>
<td>Shh-Cre;Emc3fl/fl</td>
<td>21.5% (32)</td>
<td>23.5% (35)</td>
</tr>
</tbody>
</table>

The genotypes and numbers of progeny generated are listed. Of 80 pups genotyped at weaning, no Shh-Cre;Emc3fl/fl mice were observed. For progeny analyzed at E18.5, the percentage of each genotype was consistent with the expected Mendelian ratio.
Emc3 deficiency leads to decreased ABCA3 and abnormal processing and trafficking of surfactant-associated proteins. Deleterious mutations in SFTPB or ABCA3 impair LB formation and surfactant production in both humans and mice (1, 17, 18). We therefore assessed components of the surfactant system required for normal postnatal respiration. While the expression of assessed components of the surfactant system required for normal surfactant production in both humans and mice (1, 17, 18). We therefore investigated whether EMC interacts with ABCA3 to influence its stability. To model AT2 cells in vivo, we utilized MLE-15 cells (mouse lung epithelial cell line transformed with SV40 large T-antigen) that express both ABCA3 and the surfactant proteins SP-B and SP-C (19). Consistent with its function as a subunit of the EMC complex, endogenous EMC3 was localized in the ER, as revealed by its colocalization with the ER marker, protein disulfide isomerase (PDI) (Supplemental Figure 4A). Expression of Flag-tagged ABCA3 recruited endogenous EMC3 to intracellular compartments positive for LAMP-1, indicating the interaction between EMC3 and ABCA3. A marked reduction in ABCA3 protein levels was observed in the peripheral lung of Emc3-cKO mice as assessed by immunostaining (Figure 3, A and B), immunoblotting (Figure 3G), and immunogold ultrastructural labeling (Supplemental Figure 3, A and B). The precursor of SP-B protein (proSP-B) accumulated and was abnormally secreted, while the active 8-kDa SP-B peptide (mSP-B) was nearly absent (Figure 3, A, B, and H), demonstrating a lack of proteolytic pro-

**Table 2. Quantification of the size of lamellar body–like inclusions in E18.5 AT2 cells**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Number of surveyed LBs</th>
<th>Area (mean ± SEM; μm²)</th>
<th>Maximal diameter (mean ± SEM; μm)</th>
<th>Minimal diameter (mean ± SEM; μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>377</td>
<td>0.67 ± 0.03</td>
<td>1.04 ± 0.02</td>
<td>0.75 ± 0.01</td>
</tr>
<tr>
<td>Emc3 cKO</td>
<td>388</td>
<td>0.28 ± 0.01⁴</td>
<td>0.79 ± 0.02⁴</td>
<td>0.53 ± 0.01⁴</td>
</tr>
</tbody>
</table>

Lamellar body (LB) sizes were quantified from random fields of electron micrographs of AT2 cells from control and Emc3-cKO lungs as shown in Figure 2, G and H (n = 2 each). Morphometric analysis was used to determine the interaction between EMC3 and ABCA3. A marked reduction in ABCA3 protein levels was observed in the peripheral lung of Emc3-cKO mice as assessed by immunostaining (Figure 3, A and B), immunoblotting (Figure 3G), and immunogold ultrastructural labeling (Supplemental Figure 3, A and B). The precursor of SP-B protein (proSP-B) accumulated and was abnormally secreted, while the active 8-kDa SP-B peptide (mSP-B) was nearly absent (Figure 3, A, B, and H), demonstrating a lack of proteolytic processing and misrouting of proSP-B. ProSP-C-staining was increased in the cytoplasm of AT2 cells, while the mature 3-kDa SP-C peptide (mSP-C) was abnormally colocalized with proSP-C and was not secreted into the alveolar spaces (Figure 3, C, D, and H). To determine intracellular distributions of proSP-B and proSP-C, we performed coimmunofluorescence staining with LAMP-1, which stains lysosome-related organelles, including LBs, multivesicular bodies (MVBs), and lysosomes. ProSP-B and proSP-C were colocalized with LAMP-1 in AT2 cells of Emc3-cKO mice (Figure 3, E and F), indicating retention of unprocessed precursors. Consistent with these findings, immunogold-labeled proSP-B and proSP-C were increased in the MVBs of Emc3-deficient AT2 cells (Supplemental Figure 3, C-F). Altogether, our data demonstrate that EMC3 is required for stabilization of the ABCA3 protein and for the normal processing and trafficking of SP-B and SP-C, processes that are necessary for surfactant function.

**Figure 2. Surfactant deficiency and neonatal respiratory failure in Emc3-cKO mice.** (A and B) Lungs from control and Emc3-cKO mice (P0) are shown, demonstrating diffuse atelectasis after deletion of Emc3. (C and D) Hematoxylin/eosin–stained sections show normal lung morphogenesis but collapsed airspaces in Emc3-cKO mice at P0. Scale bars: 100 μm. (E and F) Hematoxylin/eosin-stained sections show normal lung histology in Emc3-cKO mice at E18.5, the day before birth. Scale bars: 100 μm. (G and H) Electron microscopic images identify lamellar body–like inclusions (arrows) in control (G) and Emc3-deleted (H) AT2 cells at E18.5. AT2 cells from Emc3-cKO mice lacked normal lamellar bodies and contained abnormal, smaller, and heterogeneous lamellar body–like inclusions. Scale bars: 2 μm.
ing EMC3 was produced with an N-terminal c-Myc tag. When transfected into MLE-15 cells, Myc-EMC3 acted the same way as endogenous EMC3; it was readily detected in the ER, where it was colocalized with calnexin (Supplemental Figure 5A). In MLE-15 cells coexpressing Myc-EMC3 and ABCA3-Flag, the 2 proteins were primarily colocalized with LAMP-1, but not with calnexin, in intracellular compartments (Figure 4, A and B). Furthermore, Myc-EMC3 and ABCA3-Flag coimmunoprecipitated (co-IP) with other endogenous EMC subunits, including EMC1 and EMC4 (Figure 4C and Supplemental Figure 4D), indicating that ABCA3 and EMC proteins formed a complex in MLE-15 cells. RNAi-mediated knockdown of Emc1, Emc3, or Emc4 destabilized the complex (Figure 4D), demonstrating that they function together. Consistent with Emc3-cKO defects, efficient knockdown of the EMC complex by Emc3 siRNA and Emc4 siRNA decreased the stability of ABCA3-Flag in MLE-15 cells (Figure 4D). Taken together, these data support the concept that EMC3 interacts with ABCA3 to regulate its biogenesis. With similar assays, we were not able to detect direct interactions between proSP-C and Myc-EMC3 (Supplemental Figure 5, B and C), implying a distinct mechanism underlying the control of processing and trafficking of this surfactant protein by EMC3. Consistent with this finding, changes in the profiles from whole-lung mRNA in Emc3-cKO mice were distinct from those previously shown in Abca3-deficient lung (18) (Supplemental Table 1), supporting an independent role of EMC3 in surfactant biogenesis.

Figure 3. EMC3 is required for normal processing and trafficking of surfactant-associated proteins. (A and B) E18.5 control and Emc3-cKO lung sections were stained for ABCA3 and proSP-B. ABCA3 staining was reduced and proSP-B increased in AT2 cells from Emc3-cKO mice. ProSP-B was abnormally present in peripheral lung saccules in Emc3-cKO mice, indicating its abnormal trafficking. (C and D) E18.5 control and Emc3-cKO lung sections were stained for proSP-C and mature SP-C (mSP-C). Both proSP-C and mSP-C accumulated in the cytoplasm and mSP-C was not secreted from Emc3-cKO AT2 cells. (E and F) E18.5 control and Emc3-cKO lung sections were stained for proSP-B, proSP-C, and LAMP-1, a marker for lysosome-related organelles. Intracellularly retained proSP-B and proSP-C colocalized with LAMP-1 in Emc3-cKO AT2 cells. (G) Western blotting of whole-lung lysates demonstrated decreased ABCA3 in Emc3-cKO lung. (H) Western blotting of whole-lung lysates showing the accumulation of proSP-B and proSP-C, and lack of mature SP-B (mSP-B) in Emc3-cKO lung. Scale bars: 50 μm.
critical for PC synthesis cause severe and acute lung diseases mediated by AT2 cell injury or surfactant deficiency (1). Misfolding and misprocessing of surfactant proteins cause AT2 cell injury, related in part to the activation of UPR due to the misfolding and accumulation of the abnormal proteins within AT2 cells (1, 2, 14–16). Expression of disease-associated mutant ABCA3 or SP-C proteins in an AT2-like cell line modestly induced expression of EMC3 and EMC4, concomitantly with ER stress (Supplemental Figure 6).

To further interrogate the function of EMC3 in the regulation of pulmonary surfactant protein and lipid homeostasis, we isolated epithelial cells (EpCAM+ cells) from E18.5 control and Emc3-cKO lungs by magnetic cell sorting (Supplemental Figure 7). EpCAM+ cells were then subjected to RNA sequencing, proteomic, and lipidomic analyses. Data from these experiments were integrated and correlated to enable a comprehensive view and quantitative measurements of lipid and protein in AT2 cells, seeking to identify mechanisms by which EMC3 regulates AT2 cell functions. Transcriptional and/or posttranscriptional changes in Emc3-deficient epithelial cells were revealed by integration of transcriptomic and proteomic analyses. Consistent with the presence of misfolded/misprocessed ABCA3 and surfactant proteins SP-B and SP-C, both proteomic and transcriptomic analyses of Emc3-cKO AT2 cells predicted activation of the UPR signaling pathway (Figure 5, A and B, and Supplemental Table 2). The UPR comprises 3 parallel sensing pathways: IRE1α/XBP1, ATF6, and PERK/eIF2α. The PERK branch of the UPR was activated in Emc3-cKO AT2 cells, as indicated by increased expression of ATF4 (Figure 5, E, F, and I), the major effector of the integrated stress response (ISR) (20), and its downstream target, GADD34 (Figure 5I). In response to PERK-mediated phosphorylation of eIF2α, ATF4 dimerizes with other transcription factors, including ATF3 (Figure 5, C and D) and DDIT3/CHOP, to restore cellular homeostasis or, in the face of prolonged stress, to induce cell death. Likewise, the IRE1α/XBP1 branch of the UPR was activated by Emc3 deficiency. As a surrogate of IRE1α activity, splicing of Xbp1 mRNA was increased, as indicated by the increased ratio of spliced Xbp1(S) to full-length Xbp1(U) (Figure 5J). The functionally active transcription factor Xbp1(S) upregulates a series of chaperones, including HSPAS/BiP and DNAJB9/Erdj4 (Figure 5A and Supplemental Table 2), which are involved in folding of surfactant proteins (21). In contrast to the above 2 branches, ATF6 was not activated, as both the full-length ATF6(FP0) and the cleaved transcription factor ATF6(FPS0) were reduced in Emc3-cKO AT2 cells (Figure 5, G–I). Taken together, our data demonstrated that Emc3 deletion selectively induced UPR pathways in AT2 cells of the murine lung (Figure 5K).

Emc3 deletion disrupts surfactant phospholipid and protein synthesis in AT2 cells. Consistent with accumulation of surfactant proproteins (Figure 3), proteomic analysis of Emc3-cKO AT2
Figure 5. Loss of Emc3 induced the unfolded protein response in EpCAM+ sorted epithelial cells. (A and B) Heatmap of the mRNAs (A, blue/red) and proteins (B, blue/yellow) involved in the UPR pathway are shown. Proteomic and RNA sequencing data were obtained from EpCAM+ sorted epithelial cells from control and Emc3-cKO mice at E18.5. Genes and proteins were categorized by ToppGene. P values and fold changes for each mRNA and protein are listed in Supplemental Table 2. (C–H) Immunohistochemical staining for ATF3 (C and D), ATF4 (E and F), and ATF6 (G and H) was performed on lung sections from E18.5 control and Emc3-cKO embryos. ATF3 and ATF4 staining was increased and ATF6 staining was unaltered in the mutant lungs. Scale bars: 100 μm. (I) Western blots using EpCAM+ cell lysates from control and mutant lungs at E18.5 were performed using the indicated antibodies. (J) Increased Xbp1 splicing in Emc3-cKO mice. Levels of the spliced Xbp1 transcript [Xbp1(S)] were normalized to that of the full-length Xbp1 by qPCR. mRNAs were isolated from E18.5 control and Emc3-cKO EpCAM+ cells. Data are the mean ± SEM. *P < 0.05 using unpaired, 2-tailed Student’s t test. n = 4/group. (K) Model for the induction of UPR in Emc3-cKO AT2 cells. The model was built based on the integration of RNA sequencing and proteomic data. Relationships between differentially expressed genes and proteins were determined by Genomatix Pathway System (GePS) and Ingenuity Pathway Analysis (IPA) suites. System models were created using IPA’s Path Designer.
cells demonstrated a reduction of a number of proteases critical for the processing of surfactant proteins, including cathepsin H (CTSH) and napsin (NAPSA) that occurred without changes in their encoding RNAs (Figure 6, A and B). Similarly, levels of other cathepsins, CTSB, CTSC, and CTSZ, were significantly decreased (Figure 6B), which together may reflect altered sorting/trafficking of surfactant proteins to the lysosome and LB compartments.

Besides ABCA3 and UPR components, major groups of proteins were affected at RNA and/or protein levels, including enzymes and transporters involved in lipid homeostasis (Figure 6, A and B, and Supplemental Table 2). Similar to ABCA3 and other EMC subunits, some ER proteins (e.g., ACAT1/2 and ABCD3) were decreased in spite of normal RNA concentrations, indicating that these proteins may be directly or indirectly regulated by EMC3 at the level of protein folding or stabilization. RNAs encoding a number of key enzymes (e.g., PON1, SCD1/2, and ACAT1/2) and lipid transport proteins (e.g., Fabp5, SCD2, and ABCD3), some with known roles in surfactant phospholipid synthesis, were decreased (Figure 6A and Supplemental Table 2). Expression of PON1, SCD1/2, Fabp5, and SCD2 changed coordinately at both RNA and protein levels, perhaps mediated by loss of SREBP signaling in the Emc3-cKO AT2 cells. This change could be due to the transcriptional regulation of active UPR in response to the accumulation of unfolded/misfolded proteins in Emc3-cKO AT2 cells. As a master transcription regulator of cellular lipogenesis, suppression of SREBP1/2 may mediate alterations in surfactant lipid homeostasis. As a master transcription regulator of cellular lipogenesis, suppression of SREBP1/2 may mediate alterations in surfactant lipid homeostasis (22, 23). In Emc3-cKO AT2 cells, lipid concentrations were markedly decreased (Supplemental Table 3). While loss of PG moieties is consistent with defects in Abca3-mutant lungs (18, 24), differential effects among various PC species were observed beyond those changed by ABCA3 deficiency. In contrast to findings in Abca3 gene–targeted mice (18, 24), triglycerides accumulated to high levels in Emc3-cKO AT2 cells (Supplemental Table 3). Since triglycerides serve as a source of substrate for surfactant synthesis, their accumulation may indicate the disruption of normal lipid biogenesis in the absence of EMC3.

Discussion
Since the identification of the EMC complex almost 2 decades ago, our study provides the first evidence to our knowledge that the EMC3/EMC complex is essential in mammalian development. The 3 major defects in Emc3-cKO lung include disrupted surfactant
sis. Transcriptional profiling demonstrated that Abca3 and Emc3 gene-deleted mice shared few changes in their lung mRNA profile (Supplemental Table 1). While loss of ABCA3 compromised the expression of genes related to the biosynthesis and transport of fatty acids and phospholipids, enzymes and transporters primarily involved in lipogenesis were changed at RNA and/or protein levels after deletion of Emc3 (18, 24) (Figure 6 and Supplemental Table 2). Moreover, decreased SREBP signaling, perhaps mediated in part by activation of the UPR, may contribute to the inhibition of lipid synthesis in the Emc3-deleted mice. While deletion of Abca3 decreased lung PC and PG content (18, 24), the marked increase in triglyceride in Emc3-cKO lungs indicates neutral lipid accumulation and a more general disruption of cellular lipid homeostasis.

EMC3 directly bound ABCA3 and was required for its stabilization in AT2 cells, but also disrupted proteolytic processing and intracellular transport of proSP-B and proSP-C. While EMC3 did not bind to the surfactant proteins, napsin and several cathepsins known to mediate their processing were reduced in the Emc3-deleted mice. Misprocessed surfactant proteins are usually misrouted as has been shown in cultured cells (25), isolated AT2

Figure 7. Role of EMC3 in surfactant biogenesis modeled by integration of RNA sequencing and proteomic data. Model for disrupted surfactant production in Emc3-cKO AT2 cells. The model is based on the integration of RNA sequencing and proteomic data. Relationships between differentially expressed genes and proteins were determined by Genomatix Pathway System (GePS) and Ingenuity Pathway Analysis (IPA) suites. System models were created using IPA’s Path Designer.
cells (26), and in vivo (27), consistent with the abnormal accumulation of surfactant proteins in Emc3-deleted AT2 cells. Perinatal lung maturation is associated with increasing synthesis of surfactant proteins and lipids required for adaptation to air breathing. AT2 cells are highly metabolic and express high levels of the surfactant-associated proteins and lipids, rendering them sensitive to alterations in surfactant homeostasis that cause ER stress. In the present study, deletion of Emc3 caused misprocessing and accumulation of surfactant proteins and other AT2 cell proteins, as well as accumulation of triglyceride and alterations in lipid concentrations, both impairing AT2 cell functions and probably inducing ER stress. In response to ER stress, UPR pathways are induced to increase ER folding capacity and to maintain cellular homeostasis. The induction of the UPR by the loss of EM3C and other EMC subunits has been reported in previous findings. In Saccharomyces cerevisiae, deletion of Emc1-6 activated the sensor of misfolded proteins, Ire1p, and induced Hac1p splicing and the UPR (8). In Drosophila, loss of Emc3 in the retina activated Ire1 and eIF2α phosphorylation (12). Consistent with those findings, we found that deletion of Emc3 in AT2 cells activated 2 of the 3 branches of the UPR, including Ire1 and PERK/eIF2α, as indicated by XBPI and ATF4 activities, respectively. While the transcription factor XBPI induces expression of a number of chaperones involved in protein folding and ERAD, PERK inhibits translation initiation and protein synthesis, thereby reducing protein cargoes in the ER. We did not detect the activation of ATF6, a transcription factor that induces UPR target genes, including a number of ER chaperones. During ER stress, activated UPR can interfere with SREBP signaling and thus disrupt lipid metabolism. PPARγ and PPARδ, transcriptional targets of XBPI (28-30), may induce expression of Insig-1/2, thus reducing levels of active SREBP (31, 32). Likewise, ISR activation following eIF2α phosphorylation reduced active SREBP and the transcription of target genes involved in lipid synthesis (33), providing potential mechanisms by which SREBP signaling is inhibited in the absence of Emc3.

Work in Danio (13), Caenorhabditis elegans (10, 11), and Drosophila (12) demonstrated that EMCs are critical for the biogenesis of multipass transmembrane proteins, but not for secreted proteins or type I single-pass transmembrane proteins. Consistent with these findings, our data demonstrated that Emc3 and other EMC subunits directly bound to and stabilized ABCA3, a 12-pass transmembrane protein (Figure 4). While deletion of Emc3 blocked normal processing and trafficking of the secreted protein SP-B and the single membrane-spanning protein SP-C (Figure 3), our data suggest that Emc3 does not cause a generalized disruption of protein folding, since deletion of Emc3 in developing lung epithelial cells did not perturb lung growth, morphogenesis, or differentiation. These effects of Emc3 deletion were relatively specific to the processes involved in synthesis and packaging of pulmonary surfactant. Histological, RNA, and proteomic analyses did not reveal widespread disruption of signal transduction, growth, or differentiation. We utilized Shh-Cre (34) to delete Emc3 in the ventral foregut early in the processes of branching morphogenesis. Patterning of the embryonic lung is controlled by diverse signaling pathways mediated by numerous multipass transmembrane proteins, for example, DISP1 and WLS. Preservation of normal lung growth and structure in the Emc3-cKO mice supports a relatively restricted substrate specificity of proteins regulated by the EM3C/EMC complex in AT2 cells.

In summary, our findings support the critical role of EM3C and the EMC complex that coordinates (a) folding and stability of proteins critical for surfactant function at birth, (b) the UPR in the ER, and (c) regulation of lipogenesis by AT2 cells. EM3C is a critical component of an ER membrane complex that integrates surfactant protein and lipid synthesis required for air breathing after birth.

**Methods**

*Mice.* To generate the Emc3^fl^ mice, an embryonic stem cell line containing the knockout-first Emc3 allele (Emc3^fl/fl^) was purchased from The European Conditional Mouse Mutagenesis Program (EUCOMM) (Figure 1A). After obtaining germ-line transmission, the conditional allele Emc3^fl/fl^ was generated and bred with Shh-Cre (The Jackson Laboratory, stock number 005622) for endoderm-specific deletion of Emc3 (Figure 1B). Emc3^fl^ and Emc3^fl^ alleles were genotyped using primers F (GAGAAGTCCCTGCAACTCC) and R (GTCACTGCTCTGACACCTCC) and R (GTCACTGCTCTGACACCTCC). For timed matings, E.05 was determined by the presence of a copulation plug in the morning. Pregnant dams were sacrificed by CO2 inhalation and embryos were harvested at E18.5. P0 newborn pups were sacrificed by anesthesia using a mixture of ketamine, acepromazine, and xylazine and exsanguinated by severing the inferior vena cava and descending aorta. All experiments were performed using at least 4 animals of each genotype per group.

*Lung immunohistochemistry, immunofluorescence, and biochemistry.* Lungs from E18.5 embryos and P0 newborns were fixed by 4% paraformaldehyde followed by paraffin embedding. Histological staining, immunohistochemistry, and immunofluorescence were performed on 5-μm paraffin sections. Bright-field images were obtained using a Zeiss Axio ImagerA2 microscope equipped with AxioVision Software. Fluorescence images were obtained using a Nikon A1Rs inverted laser confocal microscope. For the detection of surfactant proteins in embryonic lungs by Western blots, frozen lung lobes were homogenized in phosphate-buffered saline (PBS) supplemented with Protease Inhibitor Cocktail and Protease Inhibitor Cocktail. For the detection of active SREBP and the transcription of target genes involved in lipid synthesis (33), providing potential mechanisms by which SREBP signaling is inhibited in the absence of Emc3.

**Lung immunohistochemistry, immunofluorescence, and biochemistry.* Lungs from E18.5 embryos and P0 newborns were fixed by 4% paraformaldehyde followed by paraffin embedding. Histological staining, immunohistochemistry, and immunofluorescence were performed on 5-μm paraffin sections. Bright-field images were obtained using a Zeiss Axio ImagerA2 microscope equipped with AxioVision Software. Fluorescence images were obtained using a Nikon A1Rs inverted laser confocal microscope. For the detection of surfactant proteins in embryonic lungs by Western blots, frozen lung lobes were homogenized in phosphate-buffered saline (PBS) supplemented with Protease Inhibitor Cocktail and Protease Inhibitor Cocktail. For the detection of factors involved in the UPR, EpCam^+^ sorted epithelial cells were lysed in RIPA buffer (Thermo Fisher Scientific) supplemented with the Protease Inhibitor Cocktail; for the detection of factors involved in the UPR, EpCam^+^ sorted epithelial cells were lysed in RIPA buffer (Thermo Fisher Scientific) supplemented with the Protease Inhibitor Cocktail. The following antibodies were used in this work: rabbit anti-proSP-C (WRAB-9337, Seven Hills Bioreagents); guinea pig anti-proSP-C (generated in the Whitsett laboratory, raised against the N terminus of proSP-C); rabbit anti-msp-C (WRAB-76694, Seven Hills Bioreagents); rabbit anti-proSP-B (WRAB-55527, Seven Hills Bioreagents); rabbit anti-msp-B (WRAB-48604, Seven Hills Bioreagents); guinea pig anti-ABCA3 (generated in the Whitsett laboratory, raised against the first loop of ABCA3) (35); rabbit anti-ABCA3 (WRAB-70565, Seven Hills Bioreagents); rat anti-mouse LAMP-1 (1D4B, Developmental Studies Hybridoma Bank); rabbit anti-GAPDH (G5455, Sigma-Aldrich); mouse anti-β-actin (A5441, Sigma-Aldrich); rabbit anti-Hopx (sc-30216, Santa Cruz Biotechnology); goat anti-AGER (AF1145, R&D Systems); rabbit anti-Flag (F7425, Sigma-Aldrich); mouse anti-Flag (MA1-91878, Thermo Fisher Scientific); mouse anti-c-Myc (sc-40, Santa Cruz Biotechnology); chicken anti-c-Myc (A-21281, Molecular Probes); rabbit anti-calnexin (SPA-860D 61813, StressGen); mouse anti-PDI (MA3-019, Thermo Fisher Scientific); rabbit...
anti-EMC1 (PA5-23732, Thermo Fisher Scientific); mouse anti-EMC3/TFEM111 (sc-365903, Santa Cruz Biotechnology); rabbit anti-EMC3/TFEM111 (HPA042372, Sigma-Aldrich); rabbit anti-EMC4/TFEM85 (ab184162, Abcam); mouse anti-EMC7 (sc-544440, Santa Cruz Biotechnology); rabbit anti-EMC10 (ab181209, Abcam); hamster anti-podoplanin (MA5-16113, Developmental Studies Hybridoma Bank); rabbit anti-GRP78/Bip (G9043, Sigma-Aldrich); rabbit anti-NKX2.1 (also known as TTF-1; WRAB-1231, Seven Hills Bioreagents); mouse anti-ATF3 (ab191513, Abcam); rabbit anti-ATF4 (ab184909, Abcam); mouse anti-ATF6 (70B1413, Novus); mouse anti-CHOP (2895, Cell Signaling Technology); rabbit anti-GADD34 (PAI-139, Thermo Fisher Scientific); and mouse anti-GFP (sc-9996, Santa Cruz Biotechnology).

Electron microscopy and immunogold labeling. Embryonic mouse lung tissues at E18.5 were fixed and processed using a protocol modified from a previous publication (36). E18.5 mouse lungs were fixed in situ with 2% paraformaldehyde (Electode Microscopy Sciences, EMS), 2% glutaraldehyde (EMS), 0.1% calcium chloride (Sigma-Aldrich), and 0.1 M sodium cacodylate buffer, pH 7.3 (SCB; EMS) overnight at 4°C. Fixed lungs were cut into 1- to 2-mm blocks, postfixed with 1% osmium tetroxide (EMS) and 1.5% potassium ferrocyanide (Sigma-Aldrich) in 0.1 M SCB, pH 7.3, dehydrated in a graded series of alcohol, and infiltrated and embedded with EMBed 812 (EMS). Immunogold labeling was performed as previously described (37). Electron micrographs of lung sections were acquired using a Hitachi TEM H-7650 (Hitachi High Technologies America) with an AMT CCD camera (Advanced Microscopy Techniques).

Morphometric analysis of LBs. To determine whether loss of EMC3 influences LB formation, 40 AT2 cells obtained per animal at E18.5 were randomly selected (n = 2 for control and Emc3-cKO mice) for ultrastructural morphometric analysis. Electron micrographs of AT2 cells were acquired at magnification of ×5,000 using a Hitachi H-7650 electron microscope and an AMT TEM camera, followed by manual tracings of LBs in surveyed cells using FIJI, a variant of ImageJ image analysis software. Surface areas and maximal and minimal diameters of traced LBs were estimated. Differences in surface area and diameter between control and Emc3-cKO animals were compared using the Kruskal-Wallis test, where a P value less than 0.05 was considered statistically significant.

 Constructs. The C-terminal Myc-DDK-tagged EMC3 construct was purchased from ORIGENE (catalog MR203415). To make the N-terminal Myc-tagged EMC3, Emc3 cDNA was amplified by PCR using primers F (ttgtgacccatggaacaaaaactttatttctgaagaagatctgGCAG-Emc3) and R (ttgtgacGCTAGCTGCTGTAGTTCCA) and then subcloned into the pShuttle vector (GACCAGAGCTGCTGCTTGACTC) and R (ttgtgacacTTAGAATATGGCAG-Emc3) N-terminal Myc-tagged EMC3, n = 2 for control and were randomly selected (influences LB formation, 40 AT2 cells obtained per animal at E18.5 statistically significant.

The constructs encoding hABCA3-GFP, hABCA3(E292V)-GFP, and human SFTPC variants were generated by the Weaver laboratory. ABCA3 was generated by the Whitsett laboratory. Those encoding hABCA3(L101P)-GFP were provided by Jennifer A. Wambach and Francis S. Cole (Washington University, St. Louis, Missouri, USA).

The following Silencer Select siRNAs (Thermo Fisher Scientific) were used for transfection: control siRNA (catalog 4457289), Emc1 siRNA 1 (catalog s106663), Emc1 siRNA 2 (s106664), Emc3 siRNA 1 (catalog s82552), Emc3 siRNA 2 (catalog s25553), Emc4 siRNA 1 (catalog s86111), and Emc4 siRNA 2 (catalog s86112).

Isolation of epithelial cells by magnetic cell sorting. To isolate pulmonary epithelial cells, E18.5 lung lobes were harvested from control and Emc3-cKO embryos and digested with Dispase (Corning) at 37°C for 15 minutes. The lobes were transferred to 5 ml of DMEM containing 25 mM HEPES (Gibco) and 120 U of DNASE I (Sigma-Aldrich) and dissociated with a gentleMACS Dissociator (Miltenyi Biotec). The cell suspensions were passed through a 40-μm nylon mesh strainer and then pelleted by centrifugation at 300–400 g for 10 minutes. Cell pellets were resuspended in 90 μl of autoMACS Running Buffer (Miltenyi Biotec) and 10 μl of CD326 (EpCAM) MicroBeads (Miltenyi Biotec). After incubation for 15 minutes at 4°C, 900 μl of autoMACS Running Buffer was added and cells pelleted. After washing twice with autoMACS Running Buffer, cells were resuspended in 500 μl of Running Buffer and passed through a 40-μm nylon filter cap into a round-bottom tube prior to sorting using an autoMACS Pro Separator (Miltenyi Biotec). EpCAM-positive (epithelial) and -negative (non-epithelial) cells were isolated. Cells were counted and approximately 5 × 10⁴ EpCAM⁺ cells were centrifuged and stained with NKX2.1/proSP-C antibodies and DAPI. Sorting efficiency was determined by the ratio of NKX2.1⁺ cells (epithelial) and proSP-C⁺ cells (AT2) in all DAPI-stained cells. Results from 4 control and 3 Emc3-cKO mice were quantified using at least 3 independent images per sample.

RNA analyses. Total RNA was isolated from EpCAM⁺ cells or frozen lung lobes using an RNeasy Micro Kit (Qiagen), and reverse transcription reactions were performed using an iScript cDNA synthesis kit (Bio-Rad) following the manufacturers’ instructions. Quantitative PCR (qPCR) was performed using a StepOnePlus Real-Time PCR System and TaqMan gene expression assays (Applied Biosystems). Emc3 RNA was detected using Taqman probe Mm01184718_g1 and normalized to the levels of 18S RNA (Taqman probe 4352930, Applied Biosystems). The levels of the spliced form of XBP1 (Taqman probe Mm03464496_m1; Applied Biosystems) was normalized to that of the unspliced XBP1 (Taqman probe Mm03464497_s1; Applied Biosystems). Relative expression was calculated using the ΔΔCt method and statistical significance was determined by 2-tailed unpaired Student’s t test.

RNA, protein, and lipid analyses. RNA sequencing was performed on whole-lung RNAs or RNAs extracted from E18.5 lung EpCAM⁺ mice.
cells. Sequencing and alignment were done by Cincinnati Children’s Hospital Medical Center’s Gene Expression Core. Emc3-cKO and control samples \((n = 2\) each) were analyzed. Raw reads were aligned using Bioconductor’s GenomicAlignment (38). DeSeq was used to analyze the raw gene counts and to calculate differentially expressed genes (39). AltAnalyzer was also used to determine significance (40). Genes that passed only 1 differentially expressed analysis were manually reviewed to determine relevance. All differentially expressed genes had a fold change greater than 1.2, an nbinoMTest \(P\) value less than 0.05, and reads per kilobase of transcript per million mapped reads (RPKM s) greater than 1 for both samples in at least one condition (39). The complete data set has been submitted to the NCBI’s Gene Expression Omnibus (GEO) database with the accession number GSE90969.

Mass spectrometry-based proteomic and lipidomic analyses were performed at Pacific Northwest National Laboratory on EpCAM+ cells isolated from control and Emc3-cKO fetal lungs \((n = 4\) each) at E18.5. Proteomic analyses were performed using tandem mass spectrometry (LC-MS/MS) to profile protein changes employing a label-free relative quantification approach (41). Lipid analyses were performed using LC-MS/MS employing a label-free relative quantification approach (42–44). Data were log2 transformed and median normalized within each sample. Statistically significant changes were determined using 2-tailed, homoscedastic \(t\) tests. All significantly changed proteins and lipid species had a \(P\) value less than 0.05 and a fold change greater than 1.2. The lipidomic and proteomic data were deposited and are freely available at the MassIVE data repository with the MassIVE ID of MSV000080389 (https://massive.ucsd.edu/ProteoSAFe/static/massive.jsp).

RNAs, proteins, and lipids that were significantly altered were further analyzed using functional genomics tools. ToppGene (https://toppgene.cchmc.org/) was used to detect functional enrichment of significantly altered RNAs and proteins (45). Genomatix Pathway System (GePS) (www.genomatix.de) and Ingenuity Pathway Analysis (IPA) suites (www.ingenuity.com) were used to determine relationships between differentially expressed genes and proteins based on literature mining. Predicted relationships were then manually reviewed to ensure relevance before being represented in the figures. BRB-ArrayTools was used to generate heatmaps of significant gene and protein changes (46). System models were created using IPA’s Path Designer.

Statistics. For qPCR results and quantification of Western blots, values are expressed as the mean \(\pm\) SEM. Data were analyzed using a 2-tailed, unpaired Student’s \(t\) test. \(P\) values of 0.05 or less were considered statistically significant. Statistical analyses of RNA, protein, and lipid are described in corresponding sections in the Methods.

Study approval. Mice were housed in pathogen-free conditions according to the protocols approved by the Institutional Animal Care and Use Committee at Cincinnati Children’s Hospital Research Foundation.

Author contributions. XT and MH generated the conditional allele of Emc3. XT, XL, and JAW designed experiments and analyzed the data. XT characterized the defects and performed all cell culture experiments. XT and WW examined the unfolded protein response in EpCAM+ lung cells by Western blots. JMS and YX analyzed the RNA sequencing, lipidomic, and proteomic data. CLN performed the electron microscopy and immuno-electron microscopy experiments. GC, JEK, EMZ, and CA performed the lipidomic and proteomic analyses. XT, TEW, XL, and JAW wrote the manuscript.

Acknowledgments. We thank Emily Martin, Karen Apseys, and Samriddha Ray for technical assistance. This work was supported by NIH grants RO1HL136722 (to J.A.W. and X.L.) and U01HL134745 and U01HL110964 (to J.A.W.), and grants from the National Natural Science Foundation of China 31571507 and 8136120382 and NIH grant RO1GM115995 (to X.L.). Portions of this work were supported by U01HL122703 (to C.A.) and performed in the W. R. Wiley Environmental Molecular Sciences Laboratory (EMLS), a Department of Energy (DOE) office of Biological and Environmental Research (BER) national user facility located at Pacific Northwest National Laboratory (PNNL).

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