Histone deacetylase 3 controls lung alveolar macrophage development and homeostasis

Yi Yao1,2,10, Queping Liu1,2,9,10, Indra Adrianto1,2,3,4, Xiaojun Wu1,2, James Glassbrook1,2,5, Namir Khalasawi1,2, Congcong Yin1,2, Qijun Yi1,2, Zheng Dong6, Frederic Geissmann7, Li Zhou1,2,5,8✉ & Qing-Sheng Mi1,2,5,8✉

Alveolar macrophages (AMs) derived from embryonic precursors seed the lung before birth and self-maintain locally throughout adulthood, but are regenerated by bone marrow (BM) under stress conditions. However, the regulation of AM development and maintenance remains poorly understood. Here, we show that histone deacetylase 3 (HDAC3) is a key epigenetic factor required for AM embryonic development, postnatal homeostasis, maturation, and regeneration from BM. Loss of HDAC3 in early embryonic development affects AM development starting at E14.5, while loss of HDAC3 after birth affects AM homeostasis and maturation. Single-cell RNA sequencing analyses reveal four distinct AM sub-clusters and a dysregulated cluster-specific pathway in the HDAC3-deficient AMs. Moreover, HDAC3-deficient AMs exhibit severe mitochondrial oxidative dysfunction and deteriorative cell death. Mechanistically, HDAC3 directly binds to Pparg enhancers, and HDAC3 deficiency impairs Pparg expression and its signaling pathway. Our findings identify HDAC3 as a key epigenetic regulator of lung AM development and homeostasis.
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volear macrophages (AMs), the resident macrophages in lung alveoli, are important for the maintenance of homeostasis in the airways and are involved in the development of a variety of pulmonary diseases. Recent lineage-tracing studies have demonstrated that tissue-resident macrophages (TRMs) originate from embryonic yolk sac (YS) erythro-myeloid progenitors (EMPs) and can self-renew in most adult tissues at steady state without contribution from bone marrow (BM) hematopoietic stem cells (HSCs). YS EMPs differ from mature into pMac-premacrophages (pMacs) that colonize all embryonic organs as TRMs starting at embryonic day 9.5 (E9.5). The YS EMPs also enter the fetal liver (FL) starting from E12.5, where they expand and differentiate into pMacs and/or monocytes that travel to local tissues through the blood and eventually become tissue-specific TRMs. Fetal monocytes begin to accumulate in the developing lung at E14.5 and then differentiate into immature AMs (preAMs) expressing F4/80intCD11bint, which postnatally mature into AMs expressing CD11cintSiglec-FintCD11blo,4,6,7. Like the majority of other TRMs, AMs self-maintain locally at steady state throughout adult life. Under stressed conditions, AMs are either self-maintained by local proliferation or completely replaced by bone marrow-derived cells in a phenotypic-dependent manner. A few genes have been identified as requirements for AM development and homeostasis. Granulocyte macrophage colony-stimulating factor (GM-CSF) was shown to be essential for the perinatal differentiation of fetal monocytes into preAMs and for the full maturation of AMs postnatally. Transforming growth factor-β (TGF-β) is also crucial for the embryonic differentiation of fetal monocytes into preAMs, their postnatal maturation, as well as the homeostasis of adult AMs. Both GM-CSF and TGF-β signaling pathways induce the expression of PPAR-γ, a signature transcription factor essential for the development of AMs. Macrophage identity and homeostasis require the precise regulation of gene expression that is governed by different epigenetic mechanisms, such as DNA methylation, histone modification, and chromatin structure. However, very little is known about the epigenetic factors that regulate AM development from EMPs and fetal monocytes, as well as their postnatal homeostasis.

Histone deacetylases (HDACs) are enzymes that regulate gene expression by modifying chromatin structure through the removal of acetyl groups from target histones or through direct expression by modifying chromatin structure through the homeostasis. Known about the epigenetic factors that regulate AM development, we generated Csf1rCreHdac3flo conditional knockout (cKO) mice, in which HDAC3 is deficient in the Csf1r-expressing myeloid cells, including TRMs and monocytes. Both HDAC3 mRNA and protein were efficiently depleted (>80%) in preAMs from the HDAC3cKO embryos as determined by qRT-PCR and flow cytometry, respectively (Fig. 1b, c). We next examined the fetal macrophages, preAMs, and fetal monocytes in the lung from Csf1rCreHdac3flo cKO mouse embryos and wild-type (WT) littermates between E12.5 and right after birth (P0). There was no significant alteration in the frequencies of F4/80intCD11b+ fetal macrophages and F4/80intCD11b+ monocytes in the lung between cKO and WT embryos at E12.5 (Fig. 1d, e, gates shown in Supplementary Fig. 1a). However, a modest but significant reduction in the frequency of fetal macrophages was observed in HDAC3cKO embryos at E14.5 and E16.5, followed by a gradual diminish afterwards. On the other hand, the frequency of F4/80intCD11bint preAM cells that emerged at E16.5 in the fetal lung was dramatically decreased in HDAC3cKO compared to WT littermates starting at E16.5 until birth. In contrast, fetal monocytes showed compensatorily increased distribution in the absence of HDAC3 starting at E14.5. Immunofluorescence staining of lung tissue sections at E18.5 further confirmed that the number of lung preAMs was significantly reduced in the HDAC3cKO mice (Fig. 1f). Collectively, these results suggest that HDAC3 is indispensable for the development of YS pMac-derived lung macrophages and FL monocyte-derived preAMs during embryogenesis.

Next, we investigated whether HDAC3 deficiency blocks EMP development and differentiation into pMacs and FL monocytes in the YS and FL, respectively. Interestingly, within the CD45Lin− (CD45-expressing; lineage negative) population, the frequencies of CD117intF4/80+ EMPs, CD117−F4/80+ pMacs, and CD117−F4/80+ macrophages in the YS remained unaltered in HDAC3cKO embryos compared to the WT at E9.5, E10.5, and E12.5 (Fig. 1g, h, gates shown in Supplementary Fig. 1b), suggesting that HDAC3 is dispensable for EMPs and their further differentiation into pMacs and YS macrophages. Furthermore, we observed four unique sub-populations and that a lack of HDAC3 gives rise to AM subset-specific pathway dysregulation. Moreover, loss of HDAC3 results in dysregulated mitochondrial oxidative phosphorylation and cell survival. Mechanistically, HDAC3 directly binds to Ppar gene enhancers and regulates PPAR-γ signaling during AM development. Our findings uncover HDAC3 as a key epigenetic factor in the regulation of lung AM embryonic development and maintenance after birth.
comparable frequencies of EMPs in the FL between the HDAC3cKO and WT embryos at E10.5 and E12.5, suggesting that the loss of HDAC3 also does not affect EMP seeding in the FL (Fig. 1g, h). In addition, the frequency of CD64loCD11bhi-Ly6chi FL monocytes remained unaltered during the embryonic stage (E14.5-E18.5) in the HDAC3cKO mice (Fig. 1i, j, gates shown in Supplementary Fig. 1c). HDAC3 deletion in FL monocytes, which was confirmed by qRT-PCR and flow cytometry (Supplementary Fig. 2a, b), does not disrupt their differentiation into TRMs in other organs, such as the brain, spleen, kidney, and pancreas (Supplementary Fig. 3a, b, gates shown in Supplementary Fig. 1a). Overall, HDAC3 deficiency does not affect EMPs, their ability to differentiate into pMacs in the YS, or FL monocyte development, but it may play key roles in pMac-derived macrophages and FL monocyte-derived preAMs in the lung.

HDAC3 is essential for the maintenance of AMs after birth. We next examined the AMs from the Csf1rCreHdac3fl/fl cKO adult mice. As expected, the frequency of lung AMs (CD11bhiSiglec-Fhi)
Fig. 1 HDAC3 is required for embryonic development of AMs. a qRT-PCR analysis of HDAC3 mRNA expression in lung MFs, preAMs, or AMs from C57BL/6 mice (n = 4 for postnatal day 14 (P14), n = 3 for each other time point). The expression of mRNA (b) and protein (c) of HDAC3 in preAMs determined by qRT-PCR and flow cytometry, respectively, from n = 3 Hdac3fl/fl;Cd11cCreHdac3fl/fl newborns (P0–P1). Frequencies of HDAC3-expressing (HDAC3+) AMs are shown in d. Representative flow cytometry plots: gated from CD45−CD11c+Hdac3fl/fl samples in each panel. All determined by qRT-PCR and fluorescence staining of frozen tissue sections (Fig. 2b). After birth, preAMs quickly downregulate their CD45−CD11c+Hdac3fl/fl expression, and protein expression (Fig. 2c, the remaining AMs in the HDAC3-deficient mice were found to highly express CD11b. This suggests that maintenance of mature AMs in the adult mice is very likely dependent on HDAC3. However, given that Csf1r.Cre-induced gene deletion occurs in the embryonic stage, the AM defects observed in the Csf1r.CreHdac3fl/fl cKO adult mice may be due to embryonic developmental dysregulation rather than a maintenance abnormality after birth.

Fig. 2 HDAC3 is essential for the maintenance of mature AMs in the adult mice. a Flow cytometry for CD11c and Siglec-F expression of lung AMs within CD45+ live cells from n = 7 Hdac3fl/fl;Csf1r.CreHdac3fl/fl adult mice. Frequencies of AMs are shown on the right. b Lung tissue sections from adult mice as in a. Scale bars, 100 μm. Cell numbers (n = 5 Hdac3fl/fl;Csf1r.CreHdac3fl/fl) are shown per tissue area (mm²). c Histogram plot for CD11b expression of lung AMs from n = 7 Hdac3fl/fl;Csf1r.CreHdac3fl/fl adult mice. Frequencies and absolute numbers of AMs are shown on the right. d Flow cytometry analysis of AMs from n = 10 Hdac3fl/fl, n = 17 Cd11c+Hdac3fl/fl adult mice. Frequencies and absolute numbers of AMs are shown on the right. e Histogram plot for CD11b expression of lung AMs from n = 11 Hdac3fl/fl, n = 15 Cd11c+Hdac3fl/fl adult mice. FMI of CD11b is shown on the right. f Flow cytometry analysis of AMs from n = 8 Hdac3fl/fl, UbcCre+Hdac3fl/fl adult mice. Frequencies and absolute numbers of AMs are shown on the right. Each dot represents one mouse and bars represent mean ± SD of biologically-independent samples in each panel. All P values obtained by Student’s two-tailed unpaired t test. Source data are provided as a Source Data file.

was markedly decreased in the HDAC3cKO adult mice (Fig. 2a, gates shown in Supplementary Fig. 1d). This reduction was further confirmed by immunofluorescence staining of frozen tissue sections (Fig. 2b). After birth, preAMs quickly downregulate their CD11b expression and become mature AMs. As shown in Fig. 2c, the remaining AMs in the HDAC3-deficient mice were found to highly express CD11b. This suggests that maintenance of mature AMs in the adult mice is very likely dependent on HDAC3. However, given that Csf1r.Cre-induced gene deletion occurs in the embryonic stage, the AM defects observed in the Csf1r.CreHdac3fl/fl cKO adult mice may be due to embryonic developmental dysregulation rather than a maintenance abnormality after birth.

To address this question, we next generated mice with a constitutive HDAC3 deficiency in AMs postnatally by crossing Hdac3fl/fl mice with Cd11cCre transgenic mice, given that Cd11c
were co-transferred with competitor BM cells from B6.SJL mice (CD45.1 biologically-independent samples. All CD45.1 and CD45.2 expression on CD11chiSiglec-Fhi lung AMs from inducible HDAC3-deletion (Supplementary Fig. 4a). Although normal distribution of AMs treated with TAM for 5 consecutive days, then lung AMs were HDAC3 deletion in the targeted cells was confirmed by qRT-PCR (Supplementary Fig. 4a). Although normal distribution of AMs was observed in UbcCreERHdac3fl/fl mice at 1 week post-TAM treatment (Supplementary Fig. 4b), lack of HDAC3 led to a significant reduction in the frequency and number of lung AMs 3–5 weeks post-treatment (Fig. 2f). Taken together, these findings suggest that HDAC3 also plays a key role in the maintenance of AM homeostasis, as well as their maturation after birth at steady state.

Regeneration of AMs from the BM depends on HDAC3. AMs self-maintain at steady state independently of circulating precursors. However, under certain inflammatory conditions or BM transplantation following lethal whole-body irradiation, BM-derived monocytes can repopulate the AM niche. We next sought to determine whether BM-derived AMs are also dependent on HDAC3. To do this, we established a BM chimeric mouse model by co-transferring BM cells from Hdac3fl/fl or Csf1rCreHdac3fl/fl cKO donors (CD45.2+) with competitor BM cells from B6.SJL WT mice (CD45.1+) into the irradiated B6.SJLhetero mice (CD45.1+CD45.2+) recipient mice (Fig. 3a). As expected, in the chimeric mice, lung AMs were radioresponsive and largely replaced by BM-derived cells from both WT donor and competitor mice (Fig. 3b). However, the BM from HDAC3cKO mice completely failed to reconstitute lung AMs compared to the BM from WT mice. These findings suggest that HDAC3 is not only important for embryonically-derived AMs, but also required for the regeneration of AMs from adult BM under stress conditions.

HDAC3 regulates AM subset-specific gene expression profile. Recent single-cell studies have shown that heart and liver TRMs are heterogeneous populations at steady state. To characterize whether lung AMs are also heterogeneous, we sorted CD11chi-Siglec-Fhi AMs from Hdac3fl/fl WT adult mice and performed single-cell RNA-sequencing (scRNA-seq) using the 10x Genomics platform. More than 2000 cells were analyzed and T-distributed stochastic neighbor embedding (t-SNE) dimensionality reduction analysis identified 4 major AM clusters (Fig. 4a).

Each AM sub-cluster has a unique gene expression pattern (Fig. 4b) with their signature genes (Fig. 4c). Of the AM clusters, cluster 1 was the most abundant and expressed macrophage-associated genes, such as Abcg1, Mrc1, and Mpeg1. Cluster 2 expressed genes such as Birc5, Top2a, Rrm2, and Stm1l, which are involved in the cell cycle and mitosis. Cluster 3 had relatively higher expression of antigen-presentation genes, such as H2-Eb1, H2-Ab1, H2-Aa, and Csf74. Cluster 4 was enriched in many genes associated with immune response, such as Tnf, Il1a, Cxcl1, Cxcl2, and Cxcl3. The cluster-specific gene expression pattern suggests that each AM cell subset may have a distinct role in AM maintenance and function. In order to determine whether HDAC3 is required for the maintenance of a specific lung AM cell subset, we next performed scRNA-seq on the sorted AM cells from Cd11cCreHdac3fl/flcKO mice and compared the AM clusters between HDAC3cKO and WT mice. Although the composition of clusters 1 and 2 remained almost unaltered in the KO, the frequency of cluster 3 was increased from 3.3% to 4.9%, while the frequency of cluster 4 was reduced from 4.0% to 2.4% in the absence of HDAC3 (Fig. 4d, e). This suggests that the loss of HDAC3 may affect the maintenance profiles.
**Fig. 4** HDAC3 regulates AM subset-specific gene expression profiles. CD11c<sup>hi</sup>Siglec-F<sup>hi</sup> lung AMs were sorted from \( n = 1 \) Hdac3<sup>fl/fl</sup>; Cd11cCre<sup>+</sup>Hdac3<sup>fl/fl</sup> adult mouse for scRNA-seq analysis. **a** t-SNE map of individual cells clustered from 2055 Hdac3<sup>fl/fl</sup> AM cells. **b** Heat map representing signature genes which were most highly expressed (top 10) in each AM cluster. **c** Representative violin plots showing signature genes expressed by individual AM subsets. **d** t-SNE maps of individual cells from the Hdac3<sup>fl/fl</sup> and Cd11cCre<sup>+</sup>Hdac3<sup>fl/fl</sup> (3067) AM cells. **e** The percentile of each AM subset within total AM population between two groups. **f** Radar maps depicting dysregulated pathways between two groups in each AM subset. Source data are provided as a Source Data file.
of AM subsets in a cluster-specific manner. In addition, we also observed certain cluster-specific transcriptomic changes between WT and cKO mice (Supplementary Fig. 5). Ingenuity pathway analysis (IPA) revealed that each AM cluster in the cKO mice exhibited distinct dysregulated pathways compared to that in their WT counterpart (Fig. 4f). In cluster 1, the key disordered pathways included phagosome maturation, unfolded protein response, mitochondria dysfunction, and BAG2 signaling pathways, which are associated with macrophage phagocytosis, polarization, survival, and stress responses. In cluster 2, the pathways related to cell cycle control were less enriched in the cKO cells. In cluster 3, several immune-responsive pathways were upregulated in the cKO cells, including E1F2 signaling, Th1/Th2 pathways, and dendritic cell maturation. In cluster 4, the pathways involved in cytokine signaling (i.e., IL-6 and IL-10 signaling) and acute phase response signaling were less enriched in the cKO cells, whereas Toll-like receptor signaling, neuroinflammation signaling pathway, and communication between innate and adaptive immune cells, were strongly enriched in the cKO cells. These data reveal that HDAC3 deficiency results in cluster-specific pathway dysregulation in AMs at steady state, which may lead to distinct function disorders in each AM cluster. Interestingly, when we analyzed transcriptomes of AM populations without clustering, we identified a total of 186 differentially expressed genes, including 59 upregulated genes and 127 downregulated genes in the HDAC3-deficient AMs compared to their WT counterparts (Supplementary Fig. 6a). IPA of these differentially expressed genes showed that several canonical pathways related to inflammation and immune responses were significantly inhibited in the HDAC3cKO AMs (Supplementary Fig. 6b). Moreover, the signaling pathways related to macrophage survival and apoptosis were also remarkably impaired, including TREM1 signaling29,30, NF-kB pathway31,32, and STAT3 pathway33,34. qRT-PCR analysis further confirmed the downregulation of the genes associated with these three pathways, such as Fli1, Mapk3, Map3k1, Trb2, Fgr2b, Tgb1, and Tgfb1 (Supplementary Fig. 6c). These data suggest that the inhibition of TREM1, NF-kB, and STAT3 signaling pathways might reduce cell survival of HDAC3cKO AMs, leading to the decreased total number of AM cells in the absence of HDAC3 as we observed at steady-state maintenance (Fig. 2d). On the other hand, scRNA-seq also revealed that PPAR-γ was remarkably reduced in HDAC3-deficient AMs (Supplementary Fig. 6a), suggesting that HDAC3 might control AM homeostasis through regulation of PPAR-γ.

**HDAC3 controls mitochondria function and survival of pre-AMs.** To further obtain mechanistic insights into the embryonic deficiency of AMs in the absence of HDAC3, we first performed bulk RNA-seq analysis of lung preAMs from WT and at E18.5. Based on the principal component analysis (PCA) of their transcriptomes, the molecular signatures showed a clear difference in preAMs between cKO and WT (Fig. 5a). A total of 9074 differentially expressed genes (P < 0.05) were identified in the preAMs between HDAC3cKO and WT, including 4612 downregulated and 4462 upregulated genes (Fig. 5b, Supplementary Data 1). Gene Ontology (GO) analysis further identified the altered biological pathways in the HDAC3cKO preAMs, including multiple metabolic pathways, the electron transport chain, cell proliferation, and cell death (Fig. 5c, Supplementary Data 2). Most of the identified pathways are consistent with previous studies in other cell types with HDAC3 deletion14,35,36.

Furthermore, gene set enrichment analysis (GSEA) identified some gene sets that were either depleted (normalized enrichment score, NES < 0, false discovery rate, FDR < 0.05) (Supplementary Fig. 7a, b) or enriched in HDAC3cKO phenotype (NES > 0, FDR < 0.05) (Supplementary Fig. 8a, b). Notably, the genes involved in oxidative phosphorylation were significantly downregulated in HDAC3-deficient preAMs (Fig. 5d). Many subunits of protein complexes I-V of the electron transport chain, the key component of the oxidative phosphorylation process, were diminished in the absence of HDAC3 (Fig. 5e, f). In addition, we observed a significant decrease in Mitotracker Orange fluorescence-labeled AMs from cKO, which reflects the reduced mitochondria membrane potential and indicates impaired mitochondria function in HDAC3-deficient lung preAMs (Fig. 5g). Moreover, GSEA demonstrated that the absence of HDAC3 led to a strong enrichment of genes involved in apoptosis signaling (Fig. 5h). Enhanced apoptosis of preAMs at E18.5 was further confirmed by flow cytometry (Fig. 5k). These results suggest that HDAC3 regulates preAM mitochondrial oxidative function and cell death, which may be related to defective embryonic AM development in HDAC3 cKO mice.**

**PPAR-γ regulates mitochondria function and survival of AMs.** A prior study has shown impaired development of fetal lung preAMs in Vav1-Cre-induced, PPAR-γ-deficient mice at E18.5, indicating that prenatal PPAR-γ is required for AM ontogeny. Using Csf1rCrePpargfl/flcKO mice, we further found a modest reduction of F4/80intCD11b+ YS pMac-derived fetal lung macrophages at E14.5 and a drastic loss of F4/80intCD11b+ fetal liver monocyte-derived preAMs at E17.5 from Csf1rCrePpargfl/flcKO mice (Fig. 7a), which were consistent with the phenotype observed in HDAC3cKO mice (Fig. 1d, e). As expected, Cdi1csiglec-fl mice were essentially eliminated in the Csf1rCrePpargfl/flcKO adult mice (Fig. 7a), which was consistent with the previous results using Vav1Cre- and Cdi1csiglec-fl mice. Moreover, PPAR-γ gene expression was detected in the lung fetal macrophages at E14.5, which was gradually upregulated in preAMs at E16.5 and E18.5, and maintained at high levels in the AMs from young and adult mice (Fig. 7b). Interestingly,
**Fig. 5 HDAC3 controls metabolism and cell death during AM embryonic development.** Bulk RNA-seq analysis of sorted preAMs from fetal lung of n = 3 Hdac3fl/fl;Csf1rCreHdac3fl/fl embryos at E18.5. a Principle component analysis (PCA) performed on transcriptome (Hdac3fl/fl, turquoise; Csf1rCreHdac3fl/fl, red). b Heat map showing differential gene expression (P < 0.05). c Gene Ontology (GO) enrichment of the downregulated and upregulated genes in b. The biological processes with the smallest adjusted P values by EASE Score (one-tailed Fisher’s exact P-value in DAVID system) with the Benjamini-Hochberg correction for multiple testing from each class are shown. d Enrichment plot of oxidative phosphorylation from Gene Set Enrichment Analysis (GSEA) of differentially expressed genes. NES, Normalized Enrichment Score. e Heat map depicting downregulated genes related to oxidative phosphorylation identified by GSEA. f qRT-PCR analysis of mRNA expression levels of oxidative phosphorylation-associated genes. n = 3 Hdac3fl/fl;Csf1rCreHdac3fl/fl. *P = 0.03 (Cox7b); **P = 4.2e−3 (Atp5f1), 1.2e−3 (Uqcrfs1); ***P = 1.0e−4 (Atp5g3), 2.7e−5 (Cox5a), 4.7e−4 (Cox5b), 5.7e−4 (Cox6d1), 1.2e−6 (Cox6d1), 2.4e−4 (Cox6c), 7.3e−4 (Nduf4d), 1.6e−4 (Nduf8b), 8.0e−4 (Nduf2b), 1.5e−5 (Sdhb), 2.2e−4 (Uqcrq). g Histogram plot for MitoTracker Orange dye (mitochondria membrane potential) staining of preAMs determined by flow cytometry. MFI median fluorescence intensity; n = 10 Hdac3fl/fl, n = 5 Csf1rCreHdac3fl/fl. h GSEA plot of apoptosis. i Heat map showing upregulated genes related to apoptosis identified by GSEA. Colors (red to blue) show expression values (high to low). j qPCR analysis of mRNA expression levels of apoptosis-associated genes. BMP2: n = 5 Hdac3fl/fl, n = 4 Csf1rCreHdac3fl/fl; other genes: n = 3 Hdac3fl/fl;Csf1rCreHdac3fl/fl. *P = 0.01 (Bmp2), 0.011 (Ij6), 0.026 (Tnfr); **P = 1.1e−3 (Cd38), ***P = 4.7e−6 (Mmp2), 1.3e−4 (Cndt1). k Flow cytometry of annexing V staining of preAMs; n = 8 Hdac3fl/fl, n = 7 Csf1rCreHdac3fl/fl. Each dot represents one embryo at E18.5 and bars represent mean ± SD of biologically-independent samples in each panel. All P values were obtained using the Student’s two-tailed unpaired t test. Source data are provided as a Source Data file.
PPAR-γ mRNA was not expressed in the TRMs of the kidney, liver, brain, and heart (Supplementary Fig. 9), suggesting that HDAC3-PPAR-γ axis is specifically required for the development of lung AMs, but not for TRMs in other tissues.

We next revisited a recently-published microarray dataset and analyzed gene expression profiles of lung proAMs from Pparg flox/flox WT and Cd11cCrePparg flox/flox cKO mice at day 2 after birth. As expected, the PPAR-γ signaling pathway in proAMs was severely impaired in the absence of PPAR-γ, which was consistent with our observations in Cd11cCreHdac3 flox/flox cKO mice (Fig. 7c). Interestingly, oxidative phosphorylation-related genes, which were severely diminished in HDAC3cKO mice, were robustly reduced in PPAR-γ-deficient mice (Fig. 7d–f). ChIP-qPCR analysis using MH-S cells further showed an enrichment of PPAR-γ binding sites in the promoter regions (within 3 kb of the transcription start sites) of oxidative phosphorylation-related genes, including Atp5l, Cox5a, Cox5b, Cox6a1, Ndufa4, Sdhb, Uqcrfs1, and Uqcrq, all of which were downregulated in the absence of PPAR-γ (Fig. 7e, f), suggesting a possible direct impact of PPAR-γ on mitochondria function-related genes in AMs (Fig. 7g). Moreover, the apoptosis-associated genes were highly enriched in PPAR-γ-deficient mouse compared to their WT counterparts, which was also consistent with our observations in HDAC3cKO mice (Fig. 7h–j). These data suggest that the PPAR-γ signaling pathway...
serves as a key pathway that contributes to the dysregulation of mitochondrial oxidative function and cell survival in the HDAC3-deficient AMs. Taken together, our study demonstrates that the HDAC3-PPAR-γ axis is indispensable for normal mitochondrial oxidative function and cell survival during AM development (Fig. 7k).

**Discussion**

The concept of TRMs has been refreshed in the past few years. Recent lineage-tracing studies have revealed that adult mouse TRMs are derived from embryonic YS EMP-derived macrophages and FL monocytes rather than from bone marrow-derived precursors, and they are self-maintained at steady state after...
birth\textsuperscript{1,24,37}. Although a few genes (e.g., Csf2, Pparγ, and Tgfb) have been identified as requirements for AM development, little information is known about the epigenetic regulators that instruct the development and maintenance of AMs. HDAC3, belonging to the class I HDAC family, plays a critical role in many biological processes, including circadian rhythm\textsuperscript{17}, intestinal homeostasis\textsuperscript{38}, and tissue thermogenesis\textsuperscript{14}. Here, we demonstrate that HDAC3 is required for embryonic development and postnatal maintenance of lung AMs at steady state. In addition, HDAC3 controls AM regeneration from BM progenitors after lethal whole-body irradiation. Our data further indicate that HDAC3 is dispensable for development of TRMs in the brain, spleen, kidney, and pancreas, suggesting that HDAC3 regulates TRM ontogeny in an organ-specific manner.

As the resident macrophages in lung alveoli, AMs are long lived with proliferative self-renewal capability, and control the maintenance of airway homeostasis through their capacity of clearing surfactant and cell debris at steady state\textsuperscript{1,39}. During respiratory infection or inflammation, AMs also function as antigen-presenting cells that possess antigen-presenting function\textsuperscript{47}. Furthermore, the disorders of cell cycle control and DNA damage response in the proliferating cluster (cluster 2) likely lead to less proliferative capacity of this cluster, which may also partially contribute to the reduced number of total AM cells in the HDAC3\textsuperscript{-/-}KO mice. Moreover, the disorders of cell cycle control and DNA damage response in the proliferating cluster (cluster 2) likely lead to less proliferative capacity of this cluster, which may also partially contribute to the reduced number of total AM cells in the HDAC3-deficient AM cells. On the other hand, loss of HDAC3 significantly affected the antigen-presenting function of cluster 3 and the immune-responsive function of cluster 4, suggesting that HDAC3 might crucially control the function of both AM clusters during immune responses. Further studies are needed to assess the role of HDAC3 in the functionality of individual AM subpopulations in the pathogenesis of lung diseases, such as asthma and cancer. Furthermore, it remains to be determined whether fetal immature preAMs have subpopulations and how HDAC3 regulates preAM subsets, if any, during embryonic development. However, given severe developmental defects on preAMs in the absence of HDAC3 during AM development. Given the importance of mitochondria function in immune metabolism and cell survival of macrophages\textsuperscript{48}, we reason that the loss of cluster 1 caused by impaired mitochondria function largely contributes to the reduced number of total AM cells in the HDAC3\textsuperscript{-/-}KO mice. Moreover, the disorders of cell cycle control and DNA damage response in the proliferating cluster (cluster 2) likely lead to less proliferative capacity of this cluster, which may also partially contribute to the reduced number of HDAC3-deficient AM cells. On the other hand, loss of HDAC3 significantly affected the antigen-presenting function of cluster 3 and the immune-responsive function of cluster 4, suggesting that HDAC3 might crucially control the function of both AM clusters during immune responses. Further studies are needed to assess the role of HDAC3 in the functionality of individual AM subpopulations in the pathogenesis of lung diseases, such as asthma and cancer. Furthermore, it remains to be determined whether fetal immature preAMs have subpopulations and how HDAC3 regulates preAM subsets, if any, during embryonic development. However, given severe developmental defects on preAMs in the absence of HDAC3, we currently could not obtain enough cells from the KO littermates for scRNA-seq analysis. Advanced technology, e.g., mass cytometry, may serve as a promising and powerful tool to address these interesting questions in the future\textsuperscript{19}.

Inhibition of HDACs is emerging as a promising approach to treat various types of malignant diseases, neurodegenerative diseases, inflammatory disorders, and cardiovascular diseases\textsuperscript{50}. Four pan HDACi have been approved by the FDA as anticancer agents for treatment of lymphoma and multiple myeloma, while many other pan HDACi are currently in clinical trials for treatment of lymphoma and solid tumors\textsuperscript{51}. HDACi have also been recently suggested in the treatment of asthma and other inflammatory cytokines and chemokines during inflammatory conditions. Together, our data suggest that individual AM subpopulations might be specialized for unique functions to regulate lung homeostasis and antiinflammatory immunity.

Interestingly, HDAC3 deficiency resulted in unproportionable changes in the number of AM sub-cluster cells, indicating a cluster-dependent effect of HDAC3 on AM homeostasis. Moreover, the transcriptomic changes and their associated pathway dysregulation induced by HDAC3 deletion in AMs also showed a cluster-specific pattern, i.e., mitochondria dysfunction in cluster 1, cell cycle control disorders in cluster 2, antigen-presenting dysfunction in cluster 3, and immune response dysregulation in cluster 4. Notably, the mitochondria dysfunction found in the abundant AM cluster (cluster 1) is consistent with our observations from bulk-RNA seq data showing impaired mitochondrial oxidative function and deteriorative cell death in the absence of HDAC3 during AM development. Given the importance of mitochondria function in immune metabolism and cell survival of macrophages\textsuperscript{48}, we reason that the loss of cluster 1 caused by impaired mitochondria function largely contributes to the reduced number of total AM cells in the HDAC3\textsuperscript{-/-}KO mice. Moreover, the disorders of cell cycle control and DNA damage response in the proliferating cluster (cluster 2) likely lead to less proliferative capacity of this cluster, which may also partially contribute to the reduced number of HDAC3-deficient AM cells. On the other hand, loss of HDAC3 significantly affected the antigen-presenting function of cluster 3 and the immune-responsive function of cluster 4, suggesting that HDAC3 might crucially control the function of both AM clusters during immune responses. Further studies are needed to assess the role of HDAC3 in the functionality of individual AM subpopulations in the pathogenesis of lung diseases, such as asthma and cancer. Furthermore, it remains to be determined whether fetal immature preAMs have subpopulations and how HDAC3 regulates preAM subsets, if any, during embryonic development. However, given severe developmental defects on preAMs in the absence of HDAC3, we currently could not obtain enough cells from the KO littermates for scRNA-seq analysis. Advanced technology, e.g., mass cytometry, may serve as a promising and powerful tool to address these interesting questions in the future\textsuperscript{19}.
inflammatory lung diseases. However, due to common side effects of these pan HDAC inhibitors, developing drugs with high selectivity for individual HDAC has drawn great attention. On the other hand, different subtypes of HDACs appear to play disparate roles in immune defense and disease progression. Understanding the underlying mechanism by which individual HDACs regulate immune cell development, maintenance, and function will help us to identify potential therapeutic treatments. In the present study, deletion of HDAC3 leads to repressed PPAR-γ expression in fetal immature AMs, as well as adult mature AMs, as confirmed by bulk RNA-seq and scRNA-seq analyses, respectively, suggesting the essential role of the HDAC3-PPAR-γ axis in the development and maintenance of AMs. In addition, our data imply that the HDAC3-PPAR-γ axis controls fetal immature AM development, at least partially through regulation of mitochondria function and cell survival. Interestingly, PPAR-γ binding sites are enriched near the genes associated with oxidative phosphorylation, suggesting that PPAR-γ might directly regulate mitochondrial oxidative function in AMs. On the other hand, given lack of PPAR-γ expression in TRMs from other organs, the HDAC3-PPAR-γ axis seems to be only required for TRM development in a lung-specific manner. It should be noted that a recent single-cell study revealed an enrichment of PPAR-γ+ macrophages in human lung adenocarcinoma lesions that likely promotes an immunosuppressive microenvironment. Thus, HDAC3 could be promising immunotherapeutic drugs to treat lung cancer patients by targeting the HDAC3-PPAR-γ axis in lung macrophage populations, which needs to be further investigated.

In conclusion, our data have provided evidence that HDAC3 serves as a key epigenetic regulator to control AM development at the embryonic stage, as well as in AM maintenance and regeneration after birth. Accumulated research has indicated that AMs are involved in the development of cancer, autoimmune, and inflammatory diseases in the airway. Thus, our findings may shed light on HDAC3 as a potential therapeutic target for AM-based intervention in cancer and autoimmune diseases.

Methods

Animals. Hdac3 fl/fl mice were provided by Scott W. Hiebert. C57BL/6 (Strain #000664), B6.SJL (Strain #002014), UbcCreER (Strain #008068), Csf1riCre fl/fl (Strain #000664), B6.SJL (Strain #002014), UbcCreER (Strain #007001), and Pparγ fl/fl (Strain #004584) mice were purchased from the Jackson Laboratory (Bar Harbour, ME). To generate myeloid-CreER Tg lines, body weight for 3 months. Donor BM cells were harvested from age-matched γCsf1riCre fl/fl adult lungs, as well as Csf1riCre fl/fl mice. The adult lungs were harvested, minced and incubated in 1 mg/ml collagenase D (Roche, Basel, Switzerland), 0.01% DNase I (Worthington Biochemical Corp., Lakewood, NJ), and 3% FBS (HyClone, San Angelo, TX) at 37 °C. Erythrocytes from all the samples were lysed for 3 min with 0.83% NH4Cl buffer. The cells from the adult tissues were isolated by the same method with some modifications. The adult lungs were harvested, minced, and incubated in PBS containing 1 mg/ml collagenase D (Roche), 0.01% DNase I (Worthington), and 3% FBS (HyClone) at 37 °C for 45 min. The erythrocytes were lysed as described above. All the cell suspensions were passed through a 70 μm cell strainer (BD Biosciences, San Jose, CA).

Flow cytometry and cell sorting. Single-cell suspensions were centrifuged at 450 × g for 7 min, resuspended in ice-cold staining buffer (1% PBS with 2% FBS), and placed in 96-well round-bottom plates. After incubation with purified anti-FcγRIII/II antibody (clone 2.4G2) at 4 °C for 15 min, cells were stained with a mixture of fluorescent surface antibodies at 4 °C for 30 min. The stained samples were further analyzed by flow cytometry or fixed with 2% formalin at 4 °C for 20 min prior to storage at 4 °C overnight. The full list of antibodies used can be found in Supplementary Table 1. Flow cytometry was performed with FACSaria® II or FACSElite® flow cytometer (BD Biosciences). Data were acquired using BD FACSDiva software version 8.0.2 (BD Biosciences) and analyzed using FlowJo 10.5.3 (BD Biosciences).

Mitochondria function and cell apoptosis assays. The MitoTracker Orange (Invitrogen/Thermo Fisher Scientific) stock solution was prepared according to the manufacturer’s instructions. Briefly, 50 μg of lyophilized MitoTracker Orange dye was dissolved in DMSO to a final concentration of 1 mM and stored at −20 °C in the dark. Cells were incubated with 50 μM MitoTracker Orange in the prewarmed RPMI 1640 medium at 37 °C with 5% CO2 for 15 min, followed by two washes with PBS. The stained cells were analyzed by flow cytometry (Ex 554 nm, Em 576 nm). Cell apoptosis was measured using the Annexin V Apoptosis Detection Kit (eBioscience) and was intraperitoneally administered at 50 μg/kg of mouse weight for five consecutive days.

Tamoxifen treatment. Tamoxifen (Sigma Aldrich, St. Louis, MO) was dissolved in corn oil (Sigma Aldrich) containing 10% (vol/vol) ethanol (Thermo Fisher Scientific) and was intraperitoneally administered at 50 mg/kg of mouse weight for five consecutive days.

Bone marrow chimeras. B6.SJL Heterozygous (CD45.1 fl/fl) mice were lethally irradiated once with 950 rads. Donor BM cells were harvested from age-matched and sex-matched CD45.2 Hdac3 fl/fl and Csf1riCre fl/flHdac3 fl/fl mice, as well as B6.SJL (CD45.1 fl/fl) mice. The BM cells from Hdac3 fl/fl or Csf1riCre fl/flHdac3 fl/fl mice were mixed with the cells from B6.SJL mice (at a 4:1 ratio) and then co-transferred to the irradiated B6.SJL recipient mice (10 × 10^6 mixed cells/mouse). The recipient mice were sacrificed 10 wks after reconstitution.

RNA extraction and quantitative real-time PCR. Total RNA was extracted from TRMs with mirCURY RNA Isolation Kit—Cell and Plant (Exiqon). The RNA was reverse-transcribed to cDNA with High Capacity cDNA Reverse Transcription Kits (Applied Biosystems, Foster City, CA). Quantitative real-time PCR (qRT-PCR) reactions were prepared using FastStart Universal SYBR Green Master (Roche, Mannheim, Germany). The reaction mixture contained 500 ng of lyophilized MitoTracker Orange dye, 100 μl of 1× binding buffer at 5 × 10^6/ml. After adding 3 μl of fluorochrome-conjugated Annexin V, the cells were incubated at RT for 13 min. The stained cells were washed once with 1× binding buffer and analyzed by flow cytometry.
Immunofluorescence. A piece of lung or liver freshly isolated from adult mice or embryos at E18.5 was immediately frozen in OCT compound for cryostat sectioning (5 µm per section). The frozen tissue sections were fixed with acetone for 15 min at −20 °C. After rinsing with PBS for 5 min, samples were blocked with anti-FcRyIII/I-Ab antibody at RT for 15 min followed by staining at 4 °C overnight with PE-labeled anti-Siglec-F (adult lung) or PE-labeled anti-F4/80 (fetal lung) antibody. The samples were then stained with 1 µg/ml DAPI at RT for 1 min and mounted with one drop of mounting medium (Thermo Fisher Scientific). Specimens were visualized on an Olympus FXX100 fluorescence microscope (Olympus Scientific Solutions Americas Corp, Waltham, MA) at ×10 magnification. At least 5 independent images were collected from each specimen. Images were collected using FSX-BSW (version 03.02.12) software (Olympus). The cell density of AMs (Siglec F+ or preAMs/TRMs (F4/80+) was measured based on cell counting in the fields using cellSens Dimensions Imaging Software version 1.15 (Olympus).

Bulk RNA sequencing. To achieve optimal library quality, cDNA synthesis and preamplification for total RNA of the sorted lung preAMs from WT or Csf1r+/−CreHdac3fl/flKO embryos at E18.5 were performed using the SMART-Seq v4 Ultra Low Input RNA Kit (Clontech, Mountain View, CA) following the manufacturer’s instructions. Briefly, cDNA was pre-amplified by a thermal cycler (Applied Biosystem) using 12–14 cycles and sheared into 150–400 bp fragments using a Bioanalyzer Pico sonicator (Diagenode, Denville, NJ). The DNA ends were repaired using End-It™ DNA End-Repair Kit (Epizyme, Cambridge, MA) after a single A was added at 3′ end of DNA fragment. The DNA ends were repaired using T4 DNA End Repair Kit (Epigen, Charlotte, NC). The repaired DNA fragments were ligated to adapters by T4 ligase (New England Biolabs, Ipswich, MA) after a single A was added at 3′ by Klenow Fragment (3′→5′Exo-) (New England Biolabs). MiniFlute Reaction Cleanup Kit (Quagen, Gaithersburg, MD) was used for DNA cleanup after each step. Illumina P5 and P7 primers with indicated adapters (Supplementary Table 3) were used for final library amplification. The 150–400 bp final product was purified on a 2% E-gel (Thermo Fisher Scientific). Sequencing was performed by the DNA sequencing core facility at University of Michigan using a 50 bp single end read setup on the Illumina Hiseq 4000 platform.

Single-cell RNA sequencing. Single-cell sequencing libraries were generated by an established protocol using the 10× Genomics Chromium Single Cell 3′ Reagent Kit (v2 Chemistry) and Chromium Single Cell Controller39. Briefly, 5000 FACs-sorted cells were loaded into each reaction for gel bead-in-emulsion (GEM) generation and cell barcoding. Reverse transcription of the GEM (GEM-RT) was performed using a Veriti 96-Well Fast Thermal Cycler, (Applied Biosystems) at 53 °C for 45 min, 85 °C for 5 min, and a 4 °C hold. cDNA amplification was performed after GEM-RT cleanup with Dynabeads MyOne Silica (Thermo Fisher Scientific) using the same thermocycler (98 °C for 3 min, 98 °C for 15 s, 67 °C for 20 s, 72 °C for 1 min, followed by a 12 cycle repeat at 72 °C for 1 min and a 4 °C hold). Amplified cDNA was cleaned up with SPRIselect Reagent Kit (Beckman Coulter, Brea, CA) that was followed by a library construction procedure, including fragmentation, end repair, adapter ligation, and library amplification. An Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA) was used for library quality control. Libraries were sequenced on an Illumina HiSeq4000 platform using a paired-end flow cell: Read 1, 26 (12–14 cycles and sheared into 150–400 bp fragments using a Bioanalyzer Pico sonicator (Diagenode, Denville, NJ). The DNA ends were repaired using End-It™ DNA End Repair Kit (Epizyme, Cambridge, MA) after a single A was added at 3′ end of DNA fragment. The DNA ends were repaired using T4 DNA End Repair Kit (Epigen, Charlotte, NC). The repaired DNA fragments were ligated to adapters by T4 ligase (Thermo Fisher Scientific). Sequencing was performed by the DNA sequencing core facility at University of Michigan using a 50 bp single end read setup on the Illumina Hiseq 4000 platform.

ChIP sequencing. Cell pellets from freshly-isolated AMs (1.38 × 10^6 and 0.77 × 10^6 for WT and cKO, respectively) of adult Hdac3fl/fl WT (n = 13) or Csf1r+/−CreHdac3fl/flKO mice (n = 26) were directly frozen at −80 °C. The samples were then submitted to EpiGentek (Farmingdale, NY) for ChIP, library preparation, and sequencing. Briefly, the cells were fixed with 1% formaldehyde and chromatin was isolated using a ChromaFlash™ Chromatin Extraction Kit. Chromatin was sheared using the Episonic2000 Sonicatin System. ChIP was performed using an antibody against HDAC3 (Abcam, ab7030, polyclonal, 2 µg/500 µl assay buffer) on the chromatin, and input DNA (without immunoprecipitation) was used as background. After adapter linking, the library was size selected (100–300 bp) and PCR amplified. Ten nanomolar of each sample library was provided for next generation sequencing on a HiSeq 4000.

ChIP-qPCR. The ChIP assay was performed on 80–90% confluent cultures using an EZ-Chip kit (Millipore, Burlington, MA, #17-371) and 1 µg of anti-RNA Polymerase II antibody (Upstate, #03-6218, 1 µg/ml), 2 µg of anti-HDAC3 antibody (Abcam, ab32369, clone Y145, 2 µg/ml), 1 µg of anti-PPARγ antibody (Cell Signaling, Danvers, MA, #2443S, clone 8118, 1 µg/ml), and 1 µg of rabbit IgG (Abcam, #ab171870, polyclonal, 1 µg/ml) per reaction. DNA-reaction enrichment was determined by normalization to an input genomic DNA. All ChIP experiments were obtained from independent chromatin preparations, and all quantitative real-time PCR reactions were performed in duplicates for each sample on each amplicon. Primers for the ChIP-qPCR are listed in Supplementary Table 2.
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**Author contributions**
Q-S.M., Y.Y., and L.Z. designed the study; Y.Y. performed majority of the experiments and drafted the manuscript; Q.L. performed most of the embryonic experiments; L.A., J.G., and X.W. analyzed sequencing data; X.W. prepared cDNA libraries for sequencing; N.K. assisted in qRT-PCR and flow cytometry work; C.Y. and Q.Y. maintained mouse colonies and performed genotyping; Q.-S.M., L.Z., Y.Y., and Q.L. analyzed the data; Z.D. supplied materials; F.G. contributed to the interpretation of the results; Q.-S.M. and L.Z. supervised the overall study and finalized the manuscript.

**Competing interests**
The authors declare no competing interests.

**Additional information**

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**Correspondence** and requests for materials should be addressed to L.Z. or Q.-S.M.

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