Aquaporin 5 regulates cigarette smoke induced emphysema by modulating barrier and immune properties of the epithelium

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Chronic obstructive pulmonary disease (COPD) causes significant morbidity and mortality. Cigarette smoke, the most common risk factor for COPD, induces airway and alveolar epithelial barrier permeability and initiates an innate immune response. Changes in abundance of aquaporin 5 (AQP5), a water channel, can affect epithelial permeability and immune response after cigarette smoke exposure. To determine how AQP5-derived epithelial barrier modulation affects epithelial immune response to cigarette smoke and development of emphysema, WT and AQP5−/− mice were exposed to cigarette smoke (CS). We measured alveolar cell counts and differentials, and assessed histology, mean-linear intercept (MLI), and surface-to-volume ratio (S/V) to determine severity of emphysema. We quantified epithelial-derived signaling proteins for neutrophil trafficking, and manipulated AQP5 levels in an alveolar epithelial cell line to determine specific effects on neutrophil transmigration after CS exposure. We assessed paracellular permeability and epithelial turnover in response to CS. In contrast to WT mice, AQP5−/− mice exposed to 6 months of CS did not demonstrate a significant increase in MLI or a significant decrease in S/V compared with air-exposed mice, conferring protection against emphysema. After subacute (4 weeks) and chronic (6 mo) CS exposure, AQP5−/− mice had fewer alveolar neutrophil but similar lung neutrophil numbers as WT mice. The presence of AQP5 in A549 cells, an alveolar epithelial cell line, was associated with increase neutrophil migration after CS exposure. Compared with CS-exposed WT mice, neutrophil ligand (CD11b) and epithelial receptor (ICAM-1) expression were reduced in CS-exposed AQP5−/− mice, as was secreted LPS-induced chemokine (LIX), an epithelial-derived neutrophil chemoattractant. CS-exposed AQP5−/− mice demonstrated decreased type I pneumocytes and increased type II pneumocytes compared with CS-exposed WT mice suggestive of enhanced epithelial repair. Absence of AQP5 protected against CS-induced emphysema with reduced epithelial permeability, neutrophil migration, and altered epithelial cell turnover which may enhance repair.

Introduction

Chronic obstructive pulmonary disease (COPD) is an irreversible disease usually occurring due to cigarette smoke exposure of variable duration and intensity. In the US, COPD accounts for more than 120,000 annual deaths,1 yet treatment of COPD is primarily limited to symptom relief and removal of environmental cofactors such as cigarette smoke.2 Pathologically, COPD is characterized by chronic bronchitis or emphysema. In chronic bronchitis, airway inflammation leads to increased mucus production and reduced mucociliary clearance, causing bronchoconstriction and airflow limitation. Emphysema is hallmark by destruction of lung parenchyma.3 Despite significant investigation, we do not fully understand pathologic mechanisms responsible for chronic bronchitis or emphysema.

As they serve as the interface with the outside, airway and alveolar epithelial barriers provide the first line of defense against inhaled cigarette smoke. Cigarette smoke (CS) increases epithelial barrier permeability.4-11 In addition, CS initiates an epithelial-derived innate immune response leading to release of pro-inflammatory mediators, recruitment of inflammatory cells, and expression of adhesion molecules.12,13 By modulating chemokine drive and adhesion molecule expression, epithelial cells regulate effector cell influx and thereby serve as an immunologic barrier. Both epithelial properties represent mechanisms for epithelial interactions: epithelial physical barrier is determined by how an epithelial cell interacts with neighboring epithelial cells, and epithelial immune barrier is determined by how an epithelial cell communicates with immune effector cells. How the two epithelial properties may relate to each other in response to CS has not been explored.
Communication between the epithelium and migrating inflammatory cells during smoke exposure may be crucial for subsequent development of pulmonary emphysema. Bronchial airway biopsies and lavage fluid from COPD patients with acute exacerbations and chronic smokers demonstrate elevated concentrations of IL-8, the primary neutrophil-recruiting chemokine in humans.\textsuperscript{19,35} Cigarette smoke extract (CSE) induced significant expression of the chemokine CXCL5 (LIX), a key murine neutrophil chemokine derived from lung epithelial cells.\textsuperscript{16} Cigarette smoke exposure altered the expression of the adhesion molecule and inflammatory cell receptor ICAM-1 in mouse lungs.\textsuperscript{3}

Multiple studies demonstrate accumulation of neutrophils and macrophages around small and large airways in patients with COPD.\textsuperscript{7,18} The functional severity of COPD has been shown to correlate with an increased presence of neutrophils and macrophages in the mucosa,\textsuperscript{19} although controversy exists regarding their specific contribution to smoke-induced pathology including emphysema.\textsuperscript{20,21} In murine models, CS induces inflammatory cell accumulation into the airways and lung parenchyma,\textsuperscript{22,23} and the causal role of macrophages and neutrophils in the pathogenesis of emphysema has been well-described.\textsuperscript{21,22} Studies investigating the role of lung epithelium on CS-induced murine emphysema have focused on cellular apoptosis,\textsuperscript{24} production of inflammatory mediators, and oxidant/antioxidant or protease/anti-protease balance.\textsuperscript{20} However, to date, there has been little evidence that the early changes in barrier function affect the epithelial immune responses.

Aquaporin 5 (AQP5) is an apical membrane water channel found in airway and alveolar epithelium positioned to respond to the luminal environment and participate in diverse epithelial responses. AQP5 polymorphisms can influence development and progression of COPD. A single nucleotide polymorphism of AQP5 was shown to be associated with COPD prevalence in a Chinese population.\textsuperscript{25} Subsequently, Hansel et al. demonstrated that polymorphisms in AQP5 were associated with rate of lung function decline among active smokers with COPD.\textsuperscript{26} Altered expression of human AQP5 in the bronchial tissue has been associated with lower lung function in the airways of subjects with COPD.\textsuperscript{26,27} AQP5 null mice and isolated epithelial cells deficient in AQP5 have enhanced barrier function. AQP5-mediated regulation of microtubule dynamics decreases paracellular permeability.\textsuperscript{28} We hypothesized that barrier enhancement in the absence of AQP5 mitigates CS-induced emphysema by two potentially related mechanisms: one, changes in the physical barrier, and two, alterations in immune response.

By exposing mice to cigarette smoke, we provide functional data that implicate the absence of epithelial AQP5 as a protective mechanism against development of emphysema, with reduced neutrophil migration to the alveolar space and altered alveolar epithelial cell turnover. Associated differences in key signaling proteins on neutrophils and epithelial cells may be crucial to the observed response. Our findings are novel, and confer protection against CS-induced murine emphysema due to the absence of AQP5 by a reduction in epithelial barrier permeability and a blunted innate immune response.

**Results**

The absence of AQP5 mitigates cigarette smoke-induced pulmonary emphysema, epithelial permeability and alveolar inflammatory cell accumulation

We exposed AQP5\textsuperscript{−/−} mice and WT littermates to chronic (6 mo) cigarette smoke to assess for emphysema as well as the alveolar inflammatory milieu. WT mice exposed to CS for 6 mo demonstrated a significant, 8% increase in the mean linear intercept (MLI) (Fig. 1A) and a significant decrease in surface area of airspace wall per unit of lung volume (S/V) (Fig. 1B) compared with age-matched WT mice exposed to air. MLI measures interalveolar septal distance and is an index of alveolar size.\textsuperscript{29,30} S/V measures septal loss and is an index for alveolar destruction.\textsuperscript{31,32} The change in MLI and S/V were typical of previous 6 mo exposures in this facility and in other published reports.\textsuperscript{24,33} In contrast to WT mice, AQP5\textsuperscript{−/−} mice exposed to CS for 6 mo did not have significant changes in MLI (Fig. 1A) or S/V (Fig. 1B) compared with age-matched air-exposed AQP5\textsuperscript{−/−} mice. Air-exposed controls from WT or AQP5\textsuperscript{−/−} had similar MLI and S/V values. The only group with a significant change in MLI and S/V were the WT CS-exposed mice, suggesting protection from emphysema in the absence of AQP5. Representative histologic lung sections stained with H&E demonstrate a qualitative increase in airspace only with CS-exposed WT mice in comparison to air exposed controls; significant airspace enlargement does not appear to occur in AQP5\textsuperscript{−/−} mice irrespective of CS exposure (Fig. 1C). Saving for some variability, 6 mo of cigarette smoke exposure has been shown to cause an 8–12% increase in MLI due to emphysema formation.\textsuperscript{29}

In addition to its water transport properties, our group has demonstrated that the c-terminus of AQP5 alters human airway epithelial permeability in vitro by altering microtubule assembly.\textsuperscript{28,34} We were interested in determining whether AQP5 also affected murine epithelial paracellular permeability. Using a previously developed assay of ex vivo tracheal permeability,\textsuperscript{34} we detected less extraluminal Evans blue dye (EBD) in trachea from air-exposed AQP5\textsuperscript{−/−} mice compared with trachea from air-exposed WT mice (Fig. 2A), but suspect this difference falls within a physiologically normal range given the absence of apparent baseline functional differences in AQP5\textsuperscript{−/−} mice compared with WT mice. With the addition of only 1 d of cigarette smoke exposure, trachea from WT mice had significantly more extraluminal EBD efflux compared with trachea from air-exposed WT mice. In contrast, CS-exposed AQP5\textsuperscript{−/−} mice did not demonstrate a significant increase in EBD tracheal efflux when compared with air-exposed AQP5\textsuperscript{−/−} mice. Next, we assessed for CS-altered permeability across airway and alveolar epithelial layers using a ratio of EBD concentration in the bronchoalveolar fluid (BAL) to EBD concentration in the lung in WT and AQP5\textsuperscript{−/−} mice after 4 weeks of air or CS exposure (Fig. 2B). Despite similar ratios in air-exposed WT and AQP5\textsuperscript{−/−} mice, CS-exposed WT mice had a significant increase in the BAL to lung EBD ratio compared with air-exposed mice, consistent with increased permeability across the airway and alveolar epithelial layers; AQP5\textsuperscript{−/−} mice were protected from this phenomenon. EBD lung concentration (−8–10
mice (Fig. 3C). We further quantified SPC using immunoblots of lung homogenate (Fig. 3D). AQP5−/− mice exposed to 6 mo of CS express higher levels of lung SPC compared with WT mice. Densitometry of SPC expressed as a ratio of SPC to actin was significantly increased in lungs from CS-exposed AQP5−/− mice. Collectively, this data suggests an increased ratio of type II to type I AEC presence in AQP5−/− mice compared with WT mice exposed to chronic cigarette smoke. We also quantified proliferation (Ki-67+ cells) among lung epithelial cells (CD326+ cells) after 6 mo of CS exposure in WT and AQP5−/− mice; a representative flow diagram demonstrating how CD326 positive and Ki-67+ positive lung cells were selected from CS-exposed and air-exposed mice is shown (Fig. S1A). Both WT and AQP5−/− CS-exposed mice had a 5-fold increase in % of Ki-67+ cells, suggesting CS induction of epithelial cell turnover. However, % Ki-67+ positivity was similar between CS-exposed WT and AQP5−/− cells among CD326+ cells (Fig. 3E). Of note, air and CS-exposed mice had a similar number of total lung cells and a similar percentage of CD326+ cells.

The absence of AQP5 blunts alveolar neutrophil accumulation after cigarette smoke

We used flow cytometry to identify and enumerate AQP5-induced differences in lung epithelial cells after chronic smoke exposure. Using total lung cells as the denominator, we found the % of CD326+ cells, a pan-epithelial cell marker, to be similar between CS-exposed strains (Fig. 3A). When we assessed the percentage of type I alveolar epithelial cells (AECs) identified by T1α,35 there were significantly fewer in CS-exposed AQP5−/− mice compared with CS-exposed WT mice; a representative flow plot is also shown (Fig. 3B). Type II AECs are known to proliferate and transition into type I AECs. Using SPC to identify type II AECs by immunofluorescence, we observed an increase in CS-exposed AQP5−/− mice compared with CS-exposed WT mice (Fig. 3C). We further quantified SPC using immunoblots of lung homogenate (Fig. 3D). AQP5−/− mice exposed to 6 mo of CS express higher levels of lung SPC compared with WT mice. Densitometry of SPC expressed as a ratio of SPC to actin was significantly increased in lungs from CS-exposed AQP5−/− mice. Collectively, this data suggests an increased ratio of type II to type I AEC presence in AQP5−/− mice compared with WT mice exposed to chronic cigarette smoke. We also quantified proliferation (Ki-67+ cells) among lung epithelial cells (CD326+) after 6 mo of CS exposure in WT and AQP5−/− mice; a representative flow diagram demonstrating how CD326 positive and Ki-67+ positive lung cells were selected from CS-exposed and air-exposed mice is shown (Fig. S1A). Both WT and AQP5−/− CS-exposed mice had a 5-fold increase in % of Ki-67+ cells, suggesting CS induction of epithelial cell turnover. However, % Ki-67+ positivity was similar between CS-exposed WT and AQP5−/− cells among CD326+ cells (Fig. 3E). Of note, air and CS-exposed mice had a similar number of total lung cells and a similar percentage of CD326+ cells.
Given the altered inflammatory milieu after chronic CS exposure, we were interested in knowing whether strain differences in BAL inflammatory cell profiles were present after a sub-acute CS exposure (4 weeks) thus preceding AQP5-dependent lung structural changes occurring after chronic CS exposure. Both groups exposed to CS had significant but similar increases in total bronchoalveolar (BAL) cells compared with their respective 4-week air-exposed controls (not shown). Compared with WT air-exposed controls, WT mice exposed to cigarette smoke for 4 weeks had a significant increase in BAL neutrophils (Fig. 4D). CS-exposed AQP5−/− mice had similar numbers of BAL neutrophils compared with air-exposed AQP5−/− mice, and significantly less than CS-exposed WT mice, a similar pattern to what we observed after 6 mo of CS. In contrast to BAL neutrophils, 4 weeks of CS exposure did not increase the number of BAL macrophages in WT mice. CS-exposed AQP5−/− mice had a significant, albeit mild, increase in BAL macrophages only when compared with WT air-exposed mice, but not AQP5−/− air-exposed mice (Fig. 4E). Therefore, alveolar macrophage differences in WT and AQP5−/− mice were dependent on the length of CS exposure. After 4 weeks of CS exposure, the number of BAL lymphocytes was not different in any of the four groups (Fig. 4E).

With persistent reduction in alveolar neutrophil abundance after sub-acute and chronic CS exposure as well as conferred protection from CS-induced structural changes in AQP5−/− mice, we assessed whether the presence of epithelial AQP5 directly impacts neutrophil migration using A549 cells, a human alveolar epithelial cell line (Fig. 4F). In the presence of an adenovirus control vector, AQP5 in A549 alveolar epithelial cells is undetectable irrespective of CS exposure (control). Using an adenoviral AQP5 overexpression system, we successfully increased AQP5 expression by immunoblot in A549 cells (+AQP5), which does not change with subsequent CS exposure. There were no differences in neutrophil migration with air exposure. After CS exposure, we observed a significant increase in PMN migration across A549 cells expressing AQP5 (+AQP5) compared with A549 without AQP5 expression (control).
The absence of AQP5 decreases signaling for neutrophil migration to the alveolar space

We have shown that the presence of AQP5 in lung epithelial cells alters barrier function and neutrophil transmigration. However, it is not clear whether the changes in migration are entirely due to AQP5 effects on the paracellular space, or if in addition to this, there are changes in epithelial signaling of neutrophils. To better study this, we investigated additional aspects of epithelial recruitment of neutrophils, including expression of key adhesion molecules and signaling proteins, as well as secretion of chemokines, in response to six months of cigarette smoke exposure.

CD11b expression is critical for migration of neutrophils and macrophages to sites of inflammation especially within the lung, where it interacts with ICAM-1 and other immune modulating receptors. CD11b expression is sensitive to changes in inflammatory conditions and immune modulators, the latter of which are known to be different between the apical and basolateral surfaces of the airway/alveolar epithelium, necessitating measurement in each lung compartment. We utilized established macrophage (F4–80+) and neutrophil (Gr-1+) antibodies to identify each cell population in BAL and lung, and then determined cell-specific CD11b expression by flow cytometry (Fig. S1B). We observed a significant reduction in CD11b expression on BAL neutrophils from 6

Figure 3. AQP5 alters CS-induced effects on type I and II pneumocyte presence. (A) Using flow cytometry, there was a similar percentage of CD326+ cells, a pan-epithelial marker, between WT and AQP5−/− mice after 6 mo of CS exposure. n = 7–10. (B) Among CD326+ cells, there was a significant reduction in the percentage of T1α positive cells, a type I AEC marker, in AQP5−/− mice exposed to 6 mo CS compared with similarly exposed WT mice. n = 8–10, p = 0.013. A histogram representing percent positivity and mean fluorescence intensity is shown. (C) AQP5−/− mice exposed to CS had increased SPC expression by immunofluorescence (IF), a type II AEC marker, compared with CS-exposed WT mice. (D) AQP5−/− mice exposed to 6 mos CS express higher levels of SPC compared with WT mice exposed to 6 mos CS in homogenized lung; densitometry was expressed as a ratio of SPC to actin. (E) Dot plot measuring the percentage of Ki-67+ cells among CD326+ cells demonstrates similar cluster and median values between WT and AQP5−/− mice after 6 mo of CS.
mo CS-exposed AQP5−/− mice compared with BAL neutrophils from 6 mo CS-exposed WT mice and to those from AQP5−/− and WT air-exposed mice (Fig. 5A). In contrast, CD11b expression was increased to similar levels on BAL macrophages from CS-exposed AQP5−/− and WT mice compared with those from air-exposed AQP5−/− and WT mice (Fig. 5B), suggesting neutrophil-specific differences in signaling in AQP5−/− mice exposed to CS.

Neutrophil alveolar abundance is determined by changes in chemokine gradient, altered migration across pulmonary endothelial and epithelial layers, or altered life span and removal from the alveolar space. To assess for differences in migration across the endothelium, we measured inflammatory cells in the lung interstitial compartment. Following bronchoalveolar lavage, we flushed the vasculature via the right ventricular outflow tract, extracted the lung, isolated cells into single cell suspension, and assessed them by flow cytometry. The total number of lung interstitial cells were similar between CS-exposed AQP5−/− and WT mice (not shown). Chronic CS-exposure did not induce differences in the percentage of lung interstitial neutrophils in either strain (Fig. 5B). CD11b expression on lung interstitial neutrophils was significantly higher after CS exposure in both strains, but not different between CS-exposed WT and AQP5−/− mice (Fig. 5B). This data demonstrates that significantly fewer neutrophils are present in the alveolar space of CS-exposed AQP5−/− mice despite a similar number of neutrophils with similar expression of CD11b in the lung interstitium compared with CS-exposed WT mice.

To determine if differences in neutrophil migration across the alveolar space of AQP5−/− or WT mice were influenced by differences in ICAM-1, a known receptor of CD11b, we measured it after chronic CS-exposure. CD11b expression on lung interstitial neutrophils was significantly higher after CS exposure in both strains, but not different between CS-exposed WT and AQP5−/− mice (Fig. 5B). This data demonstrates that significantly fewer neutrophils are present in the alveolar space of CS-exposed AQP5−/− mice despite a similar number of neutrophils with similar expression of CD11b in the lung interstitium compared with CS-exposed WT mice.

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is expressed by several different lung structural cells, so changes in abundance in the whole lung do not necessarily reflect differences in specific cell-types. WT mice exposed to CS had increased alveolar and airway epithelial cell ICAM-1 (labeled red) expression compared with WT mice exposed to air. When enumerated as a ratio of total cells (defined by labeled nuclei), ICAM-1 expression on airway and alveolar epithelial cells from WT mice was significantly increased compared with CS-exposed AQP5−/− mice, and compared with both air-exposed groups. CS-exposed AQP5−/− mice had similar ICAM-1 expression compared with air-exposed AQP5−/− mice. Collectively, this data demonstrates an AQP5-specific modulation of adhesion molecules in the context of cigarette smoke exposure that may contribute to differences in neutrophil migration.

As an additional determinant of alveolar neutrophil abundance, we measured chemokine secretion into the alveolar space focusing on cytokines secreted by epithelial cells. LPS-induced chemokine (LIX) levels were altered with CS exposure (Fig. 7A). In WT mice, there was a significant increase in LIX with CS exposure; however in AQP5−/− mice, there was a significant decrease in LIX with CS exposure, and CS-exposed AQP5−/− mice secreted significantly less LIX than their WT counterparts. Keratinocyte chemokine (KC) is secreted by epithelial cells, but also inflammatory cells including macrophages. KC did not significantly change with chronic cigarette smoke and was not different between WT and AQP5−/− at baseline for the mice sampled (Fig. 7B).

**Discussion**

AQP5−/− mice are protected from cigarette smoke-induced emphysema. We demonstrate a novel, dual modulatory role for AQP5 in murine epithelial barrier permeability and innate immunity. In the absence of AQP5, the epithelial barrier is less permeable with fewer alveolar neutrophils detected after sub-acute and chronic cigarette smoke exposure. The reduction in alveolar neutrophils may have occurred due to a combined effect of decreases in paracellular space, decreases in secretion of a potent epithelial-derived neutrophil chemokine LIX, and abrogated communication between neutrophil CD11b ligand with epithelial ICAM-1 adhesion molecule. A decrease in alveolar neutrophil migration likely contributed to protection from CS-induced emphysema development in AQP5−/− mice.

AQP5 is expressed on epithelial cells and not inflammatory cells, yet AQP5 effects on epithelial barrier function influenced communication with neutrophils during the innate immune response. There is precedence for the relationship of epithelial barrier to immune function in the gut and skin. Epithelial barrier disruption is causative in the pathogenesis of inflammatory bowel disease, and part of the pathogenesis of atopic dermatitis. Even though the lung is also at the interface with the outside world like the skin and gut, we have limited understanding of the influence of its epithelial barrier on the immune response. Historically, the alveolar epithelial barrier has been studied in the context of barrier disruption resulting from significant inflammation as in acute lung injury. In response to mild, chronic inflammation induced by smoke exposure, investigative focus has been primarily on the effects of barrier destruction on the cellular level, and the role of apoptosis in alveolar cell remodeling. Yet little attention has been paid to potential effects of early changes in barrier function on the pathogenesis of emphysema. Since we see changes in epithelial barrier function as early as after one day of cigarette smoke exposure, our data suggests that changes in epithelial barrier function predict altered immune responses to inflammatory stimuli including communication with other inflammatory cells.

Ample evidence exists to support the role of alveolar and airway neutrophils in development of emphysema in mice. We focused on epithelial AQP5 effects on alveolar neutrophil migration across the alveolar epithelial layer. (A) Using flow cytometry, CD11b mean fluorescence intensity (MFI) on BAL neutrophils was significantly reduced in CS-exposed AQP5−/− mice compared with CS-exposed WT mice and air-exposed AQP5−/− mice. n = 7–10 per group, *p < 0.05. CD11b MFI on BAL macrophages is similarly increased in CS-exposed WT and AQP5−/− mice. n = 7–10 per group, *p < 0.05. (B) Among total lung cells, the percentage of lung interstitial neutrophils (Gr-1+) is not increased in WT or AQP5−/− mice after 6 mo of CS exposure, but lung neutrophil expression of CD11b is significantly increased in WT and AQP5−/− mice exposed to chronic CS. n = 7–10 per group, *p < 0.05.
that changes in neutrophil migration occurred early and preceded the development of emphysema. In contrast, differences in alveolar macrophages were more subtle and variable based on duration of CS exposure in either strain of mice. Furthermore, alveolar lymphocytes increased to similar levels in WT and AQP5−/− mice after chronic cigarette smoke exposure, and are therefore less likely to be dependent on the presence of AQP5. Three, there were similar numbers of neutrophils in the lung interstitium in CS-exposed AQP5−/− mice compared with CS-exposed WT mice (yet still reduced in the alveolar space of CS-exposed AQP5−/− mice), suggesting impairment of neutrophil migration specifically across AQP5-deficient alveolar and airway epithelial layers. The same relationship did not exist between lung interstitial and alveolar macrophages.

To begin to dissect this interesting finding, we assessed signaling primarily in two ways: receptor-ligand expression and chemokine gradients, while recognizing that other factors including neutrophil half-life and rates of removal can be influenced by other cell types. Expression of CD11b, the ligand primarily responsible for neutrophil migration across the alveolar epithelial layer, was similar on lung interstitial neutrophils between WT and AQP5−/− mice after cigarette smoke exposure. However, neutrophils from CS-exposed AQP5−/− mice that successfully migrated across the basolateral surface to the apical surface of the alveolar epithelium expressed less CD11b compared with neutrophils from WT mice. On alveolar epithelial cells, ICAM-1 localizes to the apical surface and may regulate alveolar neutrophil abundance by tethering to it as has been shown in other mucosal surfaces. Reduced ICAM-1 expression on alveolar epithelial cells from AQP5−/− mice may also downregulate alveolar neutrophil CD11b expression, since the signal for epithelial cell – neutrophil interaction is diminished. Once the cell-cell communication is disrupted, neutrophils may be subject to enhanced apoptosis and removal as an alternative explanation for differences in alveolar neutrophil abundance after chronic CS exposure.

While ICAM-1 is integral to communication with CD11b on the apical surface, it is not thought to be expressed on the basolateral surface of alveolar epithelial cells, and therefore may not be responsible for interaction with CD11b on lung neutrophils located in the interstitium, which may explain why we did not observe differences in neutrophil CD11b expression in that compartment. We do not know how AQP5 regulates ICAM-1 expression, but one explanation may be via AQP5-directed coordination of microtubule (MT) dynamics. AQP5-derived MT changes could rearrange cytoskeleton and alter epithelial ICAM-1 expression, which others have shown to occur especially
when epithelial cells are in communication with inflammatory cells. We also observed AQP5-mediated differences in alveolar secretion of an epithelial-derived neutrophil chemokine, LIX, after cigarette smoke exposure. Whether cross-talk between the CD11b-ICAM-1 signaling pathway and the LIX chemokine gradient exists to retard neutrophil migration in AQP5-null mice is not currently known, though it is interesting that alveolar neutrophil CD11b expression and BAL LIX secretion were lower in CS-exposed AQP5−/− mice even compared with air-exposed mice, suggesting active feedback induced by CS. We cannot exclude the possibility that differences in macrophage sub-populations, their secreted products, or their interactions with epithelial cells were present between WT and AQP5−/− mice and may have affected emphysema development. Macrophages secrete proteases including MMP-2, MMP-9, MMP-12, cathepsins K, L, S, and elastase; in COPD patients compared with normal smokers, macrophages secreted more inflammatory proteins and have greater elastolytic activity. Hautamäki demonstrated the critical role for macrophage-derived elastase, a metalloproteinase that solubilizes many extracellular matrix proteins, in emphysema development after chronic CS exposure in mice. Microtubule stability was found to blunt elastase-derived epithelial cell detachment and IL-8 release, an interaction which may be altered by AQP5 expression. In addition to changes that implicate protection from lung structural damage in AQP5−/− mice, there is also evidence supporting an increase in epithelial repair. Compared with CS-exposed WT mice, CS-exposed AQP5−/− mice had an increased number of type II AECs pneumocytes and a reduced number of type I pneumocytes despite similar numbers of total and proliferating lung epithelial cells. Because the most widely accepted dogma is that type II AECs proliferate and differentiate to type 1 AECs, we postulate a decrease in type II AEC to type I AEC transition in CS-exposed AQP5−/− mice as a protective mechanism against emphysema, potentially due to type II AEC-mediated injury repair or surfactant secretion to maintain airflow and alveolar stability. AQP5 expression is confined to type I AECs; we are not aware of any known role in modulating type II AEC to type I AEC transition. However, Borok et al. have shown that there is increasing AQP5 abundance during the transition and therefore AQP5 absence could conceivably impact on the transition to full type I AEC differentiation in models of injury and repair. Human and murine studies have suggested that reduction in AQP5, a water channel, is associated with mucous hyperproduction and may help explain the pathology of airway obstruction. To our knowledge, we are the first to observe a relationship between AQP5 and development of cigarette-smoke induced structural changes consistent with emphysema. AQP5 is tightly regulated in lung epithelial cells, and dynamically responds to several pathologic stimuli including TNF-α and LPS. Acute or chronic CS exposure may also influence alveolar epithelial AQP5 expression and subsequently impact lung structure. We have previously shown that decreased AQP5 lowers levels of assembled microtubules, leading to decreased paracellular permeability. Published data suggests that microtubule disassembly induced by nocodazole decreases paracellular permeability in the lung epithelium due to altered membrane expression of E-cadherin. A reduction in airway E-cadherin expression correlated with increased barrier permeability in response to CS exposure, an effect dependent on the presence of AQP5. Coordination of microtubule dynamics with resultant changes in paracellular permeability may be an additional explanation for tight AQP5 regulation in the lung epithelium, besides well-established AQP5 regulation of transmembrane water flux. Decreased microtubule assembly due to the absence of AQP5 may affect epithelial signaling pathways which modulate epithelial immune responses. Furthermore, the absence of AQP5 may have adaptive, indirect effects on prevention of permeability and emphysema. Kneidinger showed that activation of the Wnt-B-catenin pathway, recently shown to be prominent in epithelial repair, actually protects against development of experimental emphysema. Activation of the Wnt-B-catenin pathway increases lung AQP5 levels. One could hypothesize that with a lack of negative feedback, the absence of AQP5 may lead to constitutive upregulation of the Wnt-B-catenin pathway as an adaptive response to promote epithelial repair after cigarette smoke and protect against emphysema. We present novel data demonstrating AQP5-induced alteration in epithelial barrier predisposing the epithelium to alter its innate immune response to cigarette smoke. Absence of AQP5 minimizes neutrophil migration to the alveolar space and protects against development of emphysema. Further understanding of AQP5-mediated mechanisms changing the epithelial barrier and subsequent innate immunity are necessary, and may provide insight into development of COPD.

Materials and Methods

Animal use and care
Male AQP5−/− mice (on a C57BL/6 background) and wild type (WT) littermates were bred and housed at the Johns Hopkins University Asthma and Allergy Center. Experiments were conducted under a protocol approved by the Johns Hopkins Animal Care and Use Committee.

Murine smoke exposure
WT (littermates) and AQP5−/− mice were exposed in the Johns Hopkins smoke exposure core. Mice were exposed to cigarette smoke 5 h/day, 5 d/week for 4 weeks or 6 mo by burning 3R4F reference cigarettes (2.45 mg nicotine/cigarette; Tobacco Research Institute, University of Kentucky) using a smoking machine (Model TE-10, Teague Enterprises).

Animal harvesting
Mice were anesthetized with intraperitoneal ketamine/acetylpromazine (150/13.5 mg/kg) prior to harvest. At specified time points after cigarette smoke or air exposure, 7–10 animals from various groups were anesthetized and killed by exsanguination from the inferior vena cava. The lungs were perfused free of blood with 1 ml of phosphate-buffered saline (PBS) unless otherwise specified.

Analysis of bronchoalveolar lavage (BAL)
BAL was obtained, total and differential cells counted, and BAL protein assessed as previously described. Keratinocyte-derived
Mouse trachea were excised, and both ends were cannulated with 18-gauge catheters. In one end, we instilled 50 μL of EBD-4% albumin into the lumen of the trachea for 20 min. The tracheal lumen was gently flushed with PBS, homogenized, and processed as above. The supernatant (200 μl aliquots each in 96-well microtiter plates) was collected and the absorbance of tracheal, lung, or BAL supernatants, and serum (diluted 1/10) was measured by spectrophotometer at 620 nm wavelength using a standard curve prepared with serial dilutions of formamide. EBD concentration was multiplied by the dilution factor (3 ml + wet weight of R lung) to account for PBS, formamide and water weight of the R lung. Data are presented as EBD μg/R lung, or as a ratio of BAL/Lung EBD.

**Lung histology and morphometry**

Left lungs were collected and processed for histology and morphometry by inflating to 25 cm H2O with 0.6% agarose, isolated, sectioned (5 μm), and stained with hematoxylin and formalin (H&E). Fifteen images per mouse were captured at 100X (Nikon Eclipse 50i), and mean linear intercept (MLI) and surface area of airspace wall per unit of lung volume (S/V) ratio were measured in BAL and culture medium by ELISA.

**Evans blue dye (EBD) assay**

*In vivo, alveolar.* Mice were injected with 20 mg/kg EBD (0.5% Evans blue dissolved in 4% albumin/0.9% saline (PBS, retro-orbital vein) one hour prior to harvest. At harvest, blood was collected from the aorta. The pulmonary circulation was flushed with PBS, followed by lavage and ligation of the right lung, which was then frozen (~80°C). The frozen right lung was homogenized in 1 ml PBS, and then centrifuged at 5,000 x g (7800 rpm, 30 min). The homogenate was diluted in 2 ml of formamide and incubated at 60°C for 18 h.

*Ex vivo, trachea.* Mouse trachea were excised, and both ends of the trachea were cannulated with 18-gauge catheters. In one end, we instilled 50 μL of EBD-4% albumin into the lumen of the trachea for 20 min. The tracheal lumen was gently flushed with PBS, homogenized, and processed as above. The supernatant (200 μl aliquots each in 96-well microtiter plates) was collected and the absorbance of tracheal, lung, or BAL supernatants, and serum (diluted 1/10) was measured by spectrophotometer at 620 nm wavelength using a standard curve prepared with serial dilutions of formamide. EBD concentration was multiplied by the dilution factor (3 ml + wet weight of R lung) to account for PBS, formamide and water weight of the R lung. Data are presented as EBD μg/R lung, or as a ratio of BAL/Lung EBD.

**Lung immunofluorescence**

Right lungs were isolated, sectioned onto slides, and submerged into 4% formaldehyde/10% formalin in PBS (≤ 48 h, room temp), and then heated (15 min, 60°C). Slides were deparaffinated and hydrated with xylene (2×, 1 min), 100% EtOH (2×, 1 min), 95% EtOH (2×, 1 min), 80% EtOH (1×, 1 min), and then water (1×, 1 min). Citrate buffer (fresh, salts) was added to the slides (20 min, 95°C), followed by passive cooling to room temperature. Samples were fixed with hydrophobic pen, then blocked (20% serum, 1% BSA, 0.5% Tween-20 in PBS), rinsed (0.3% Triton X-100, 1% BSA), and covered with primary antibody for E-cadherin (1:200, Cell Signaling, #3195S) or ICAM-1 (1:200, Biolegend, #116110) in PBS with 0.3% Triton X-100, 1% BSA for 18 h (4°C). We then washed (2×, 10 mins, 0.3% Triton X-100, 1% BSA), and added a secondary antibody (1:400, for E-cadherin-Ax488 Donkey Anti-rabbit IgG, for ICAM-1 Ax549 Donkey anti-mouse IgG, Invitrogen) diluted in 0.3% Triton X-100, 1% BSA (2 h, 25°C) in the dark. SPC primary antibody was diluted in 4% serum/DPBS (1:100, Santa Cruz, Cat #13979), washed (3×, 10 min, DPBS), biotinylated (1:1000 Biotin goat anti-rabbit, Invitrogen, Cat # B-2770), washed again (2×, 5 min, DPBS), and then we added a secondary antibody (1:400, Alexa Fluor 488 streptavidin antibody, Invitrogen, Cat # S-11223) diluted in 4% serum/DPBS (2 h, 25°C) in the dark. After subsequent rinse (2×, 10 min in PBS, then 1×, 10 min in TBS (pH 8.6)), slides were covered with Fluoromount, sealed, and stored at 4°C. Quantification: ImageJ software by quantifying the ratio of mean fluorescence intensity between the protein of interest and the intensity of the nuclei in the same region of interest. This was done from a minimum of 3 different animals per condition.

**In vitro smoke exposure and neutrophil migration**

A549 cells (ATCC) were grown on collagen-coated inserts at 37°C with 5% CO2 in media (450 mL F12K, 50 mL FBS). We used a Vitrocell smoke chamber to deliver tobacco to A549 cells to mimic tobacco smoke exposure of human epithelium in smokers. The cells were cultured on a 6-well insert as described above allowing medium on the bottom of the porous insert, while the top was exposed to either air (control) or cigarette smoke. We exposed cells to 2 cigarettes (7 min/cigarette). After exposure, the inserts were inverted into a larger dish. Freshly isolated peripheral blood PMNs (courtesy of Bruce Bochner) were labeled with a commercially available fluorescent probe (carboxyfluorescein diacetate (CFDA); excitation/emission spectra similar to FITC to allow PMN tracking. Media containing the labeled PMNs were applied to the basolateral membrane of the inverted insert. After 4 h, we collected the basolateral media; inserts were collected and lysed to determine sample fluorescence to objectively quantify PMN transmigration.

**Immunoblotting**

Lungs or A549 cells were harvested and lysed in RIPA buffer (50 mM TRIS-HCl pH 8, 150 mM NaCl, 1% Triton X-100 + Roche Protease Inhibitor, 15 min), followed by spin, and Bicinchoninic acid assay (BCA Pierce) for total protein in the whole cell lysate. Normalized proteins were loaded onto 10% gel. We then blocked with 4% BSA for 30 min, and incubated with SPC antibody (Santa Cruz Antibody, Cat # 13979), washed (3×) with TBST, then incubated with secondary in 4% BSA (Bio-Rad, Cat # 170-6515) for an hour. Followed by wash and ECL with HyGLO (Denville Scientific, Cat #E2400). Antibodies to the carboxy-terminus of human AQP5 were generated and probed for as before.

**Flow cytometry**

Cells were processed for flow and analyzed as previously described, using the following antibodies (or relevant isotypes): anti-LY6G-FITC, anti-CD11b-APC-Cy780, anti-F4/80-APC, anti-CD86-PacBlue, anti-CD326-APC Cy7, anti-T1o-APC, and anti-K,67-PE. Mean fluorescence intensity (MFI) was based on the geometric mean expression for a particular marker among cells gated by positive expression.
Statistical analysis
All values are reported as mean ± SEM except Figure 3E, where individual values and the median for each group are shown. Multiple groups were compared using one-way ANOVA with Bonferroni t-test for multiple pairwise comparisons when data are normally distributed or with Kruskal-Wallis assessment on ranks when data are not normally distributed. Two groups were compared using the student’s t-test or Mann-Whitney rank sum test when sample size or variance was not equal. Statistical analysis was performed using Sigmaplot 11.0 (Systat Software). A p < 0.05 was used for significance.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Supplemental Materials
Supplemental materials may be downloaded here: www.landesbioscience.com/journals/tissuebarriers/2013TISSBARRIER028R1-Sup.pdf

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