Development of an airway mucus defect in the cystic fibrosis rat

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Introduction

Cystic fibrosis (CF) is the most common autosomal recessive genetic disorder, resulting from dysfunction of the CF transmembrane conductance regulator (CFTR) protein (1). More than 1,700 disease-causing mutations in the CFTR have been identified, many of which result in little or no CFTR activity (2). CFTR transports both chloride and bicarbonate ions across the epithelial surface of many tissues (3–5); in the airway, this provides the epithelial surface with adequate hydration, osmotic pressure, and pH regulation (6–8). Deficient CFTR anion transport, and thereby reduced chloride and bicarbonate concentrations at the airway surface, leads to decreased mucus clearance, copious airway obstruction, and loss of lung function (1). Bicarbonate concentration, in particular, has been implicated as crucial to promoting appropriate mucin unfolding and normal mucus properties (9–12). Without bicarbonate, mucus release from both goblet cells and airway submucosal glands is impaired (10, 12–14). In the setting of abnormal mucus, hyperosmotic periciliary layer (PCL) depletion also contributes to mucus stasis (15, 16).

The mechanisms underlying the development and natural progression of the airway mucus defect in cystic fibrosis (CF) remain largely unclear. New animal models of CF, coupled with imaging using micro-optical coherence tomography, can lead to insights regarding these questions. The Cftr−/− (KO) rat allows for longitudinal examination of the development and progression of airway mucus abnormalities. The KO rat exhibits decreased periciliary depth, hyperacidic pH, and increased mucus solid content percentage; however, the transport rates and viscoelastic properties of the mucus are unaffected until the KO rat ages. Airway submucosal gland hypertrophy develops in the KO rat by 6 months of age. Only then does it induce increased mucus viscosity, collapse of the periciliary layer, and delayed mucociliary transport; stimulation of gland secretion potentiates this evolution. These findings could be reversed by bicarbonate repletion but not pH correction without counterion donation. These studies demonstrate that abnormal surface epithelium in CF does not cause delayed mucus transport in the absence of functional gland secretions. Furthermore, abnormal bicarbonate transport represents a specific target for restoring mucus clearance, independent of effects on periciliary collapse. Thus, mature airway secretions are required to manifest the CF defect primed by airway dehydration and bicarbonate deficiency.
KO rat), which enables experiments out to at least 6 months of age and provides a model appropriate for therapeutic intervention coupled with mechanistic studies as the disease evolves. Preliminary characterization demonstrated completely absent CFTR expression and function in all tissues, including the airways, combined with initial events that characterize the CF lung, namely a dehydrated airway surface (26). Additionally, the rat expresses a higher number of submucosal glands in the upper airways, especially tracheae and bronchi (27), a feature that has been shown to be important to human CF disease (28–33).

In the present study, we combine traditional methods with 1-micron resolution optical coherence tomography (μOCT) to show that airway submucosal glands are developmentally delayed in the CFTR KO rat but are required for the development of an abnormal mucus defect. Neither hyperacidic pH nor dehydration of the airway surface can impart mucus stasis unless gland secretions are mature, lending a crucial insight into the role of submucosal glands in the development of CF airway disease. Further, gland secretions are particularly sensitive to changes in bicarbonate concentration, but not to changes in pH, for the initiation of delayed mucus clearance. This is a finding with important therapeutic indications, as mucus clearance is an important component of host defense and effective CF treatment.

**Results**

**Submucosal gland development is delayed in the CFTR KO rat trachea.** The phenotype of CFTR KO rats at weaning was recently described (26). At 4–6 weeks of age, overall tracheal morphology appeared similar between KO and WT rats, including the number and intensity of mucus-producing cells in the trachea. As rats aged, submucosal glands in the trachea became more apparent in KO rats and demonstrated submucosal gland duct plugging by 6 months, features absent in WT rats (Figure 1A). Morphometric measurements of WT and KO tracheal histology images indicated that submucosal gland area was diminished in KO rats at weaning but became greater than WT by 6 months of age (0.172 ± 0.3 mm² WT vs. 0.244 ± 0.04 mm² KO, \(P < 0.05\), Figure 1B). Correspondingly, the area of intracellular mucus staining became larger in KO rats compared with that in WT rats by 6 months of age (Figure 1C), even though KO rats remained smaller overall compared with their WT littermates (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.97199DS1). Although male rats of both genotypes grow larger than females of the same age, there was no detectable sex-related difference in histomorphometry. To assess the function of tracheal submucosal glands and the effect of their secretions on the airway surface at 1, 3, and 6 months of age, tracheae were imaged under μOCT as previously described (22, 24, 26, 34) and the airway surface liquid (ASL) depth was measured before and after cholinergic stimulation with acetylcholine (Figure 2A). Comparison of ASL depths of KO rats to those of their age-matched WT littermates showed that ASL depths were diminished in both basal and stimulated conditions, from weaning until at least 6 months of age (WT 1 month 22.9 ± 4.6 μm, 6 months 43.6 ± 12.5 μm vs. KO 1 month 6.9 ± 0.7 μm, 6 month 19.5 ± 4.8 μm, \(P < 0.05\) at all ages, Figure 2, B and C). Compared with WT tracheae, those of KO rats had a diminished response to cholinergic-stimulated gland secretion at 1 month of age but then exceeded the WT response by 6 months of age. The absolute change in ASL depth in acetylcholine-stimulated tissue, compared with baseline, showed that WT rat tracheae responded as expected to cholinergic stimulation (28, 31, 35), with increasing ASL depths at all ages. In contrast, KO rats had an almost undetectable difference in ASL depth upon cholinergic stimulation at 1 month of age, whereas fluid secretion in response to acetylcholine was greater by 3 and 6 months (Figure 2, A and D). When analyzed as percentage change over baseline, WT rats approximately doubled the volume on the surface at 1 month of age and increased the volume by about 50% at later time points (Figure 2E). KO rats, however, exhibited a minimal volume increase at 1 month but almost doubled ASL depth at 3 and 6 months of age (KO 1 month, −1.2% ± 0.1%, 3 months 143.3% ± 41%, 6 months 95.3% ± 12.2%). These data suggest that maturation of the submucosal glands in the trachea of the KO rat is delayed, not only in size but also in function compared with their WT littermates; additionally, eventually stimulated secretion response in KO rats exceeded that of normal rats.

**Absence of CFTR alters airway surface properties of the KO rat.** KO rat tracheae exhibited characteristics of the airway that have been reported in other models of CF. The pH of the airway surface, measured by a small-volume pH probe, was hyperacidic compared with WT littermates, at all ages tested (Figure 3A). While the pH of both the WT and the KO tracheae increased slightly following cholinergic stimulation (by 0.24 ± 0.009 and pH 0.36 ± 0.007 pH units, WT and KO, respectively), the pH of KO tracheal surface remained more acidic than that of the WT tracheal surface (Figure 3B). The solid content of the mucus,
which has been reported to be increased in the CF airway (16, 36–38), was also measured in both WT and KO rat tracheae as the rats aged. The percentage of solid content of the airway mucus layer under basal conditions was increased significantly in KO rats at all ages evaluated (WT 1 month, 2.31% ± 0.3%, 3 months 2.96% ± 0.6%, 6 months 0.84% ± 0.1% vs. KO 1 month 13.31% ± 0.5%, 3 months 9.93% ± 0.6%, 6 months 10.51% ± 1.6%, P < 0.001 at all ages, Figure 3C). This result was consistent with reduced ASL depth and reflective of significant dehydration (Figure 2, A and B), indicating that the KO rat exhibits both characteristics (dehydration and abnormal pH) postulated to drive CF pathogenesis.

Nevertheless, the effects of aging and gland secretion on this compartment of the airway surface was more complicated than the initial steady-state condition. At a young age, cholinergic stimulation resulted in reduced solid content in the KO tracheae, while the 3- and 6-month-old KO rat tracheae produced an increased percentage of solid content in response to acetylcholine (Figure 3D). Examining these data as a change from baseline after cholinergic stimulation demonstrated that the 1-month KO condition is unique; this is the only age at which the solid content percentage of the mucus decreases after treatment with acetylcholine (Figure 3E). μOCT-based PCL depth measurements of the same animals showed that this layer was collapsed under basal conditions at 1 month, indicating local dehydration in KO rats compared with WT at baseline (Figure 3F), but, as with the percentage of solids, the effects of gland maturation made this more complicated. With stimulation of mucus secretion by acetylcholine treatment, PCL depths were greater in the KO than the WT rat at 1 month of age (4.46 ± 0.5 μm WT vs. 5.45 ± 4.2 μm KO, P < 0.01), no different at 3 months (4.56 ± 0.2 μm WT vs. 4.05 ± 0.4 μm KO), and ultimately demonstrated PCL collapse at 6 months of age (4.10 ± 0.2 μm WT vs. 3.53 ± 0.2 μm KO, Figure 3G). This differential response indicates that although PCL depletion is present at 1 month, gland stimulation is principally aqueous at this age, which reduces solid content and restores the local PCL depth after induction. These
Data further suggest that as airway submucosal glands develop over time, including maturation of the gland secretions themselves, increased solid content is delivered to the surface, contributing to PCL collapse, probably through increased osmotic force (16, 39) induced by elevated solid content.

Altered mucus transport and viscoelastic properties develop as the CFTR KO rat ages. Next, we evaluated the effect of changes in PCL depth, solid content, and pH induced by gland expression and stimulation on mucus transport to assess their effect on epithelial function and host defense. By 6 months of age, the KO rat expressed increased mucus secretion and accumulation of mucus in the small airways (area 0.5–1.5 mm), coupled with an increase in the size of goblet cells, despite a similar proportion of goblet cell numbers (Supplemental Figure 2); nevertheless, as opposed to larger CF animals, there was not consistent evidence of complete occlusion of the small airways with mucus or an increase in the incidence of spontaneous infection. However, detailed investigation of the mucus properties as the rat aged revealed abnormalities. The same μOCT images that yielded measurements of ASL and PCL depths were also used to calculate mucociliary transport (MCT) rates in WT and KO tracheae. At baseline, MCT rates measured in the KO tracheae were no different than those of WT littermates; however, as the rats aged, mucus transport became significantly delayed, culminating at age 6 months of age (Figure 4A). Moreover, following cholinergic stimula-

**Figure 2. Airway surface liquid depletion throughout development.** (A) Representative μOCT images of WT and KO rat tracheae at baseline conditions as well as treated with acetylcholine to stimulate mucus secretion, depicting the epithelial layer (ep), mucus layer (mu), lamina propria (lp), and airway surface liquid (ASL) depth (red bar). White bar: 10 μm (in 2 dimensions). Raw images were acquired at an 0.7 μM resolution. Quantification of the images yields measurements of ASL depth at baseline (B) and after cholinergic stimulation (C) as well as the absolute change (D) and percentage change (E) in ASL after acetylcholine stimulation. Data are shown as mean ± SEM. Data were analyzed by 2-way ANOVA. *P < 0.05, ****P < 0.0001. Data in A are representative of 3 independent studies. Data in B–E are from 6 animals/group.
tion, MCT rates were preserved in 1-month-old KO rats but were dramatically lower in the KO tracheae by even 3 months of age (Figure 4B). Particle-tracking microrheology, a robust technique that correlates highly to other methods of viscosity measurement, such as fluorescence recovery after photobleaching (24, 40, 41), was used to calculate effective viscosity of the mucus on the surface of the trachea in situ. Representative tracings indicate the Brownian motion of particles in the mucus of the defined condition and demonstrate the constraints on the movement of particles in more viscous mucus (Figure 4, E–G). Under baseline conditions at 1 month of age, effective viscosity was no different in KO tracheae compared with WT tracheae (1.76 ± 1.0 cP WT vs. 1.90 ± 1.7 cP KO, Figure 4C). At 3 months, effective viscosity of KO airway mucus increased slightly but was still no different than that of WT airway mucus (5.12 ± 1.0 cP WT vs. 10.72 ± 2.7 cP KO, Figure 4C). By 6 months of age, KO tracheal mucus was 20-fold more viscous compared with WT littermates (2.91 ± 0.9 cP WT vs. 65.09 ± 3.6 cP KO, P < 0.05, Figure 4C), consistent with delayed transport.
at this age. In the stimulated conditions, airway mucus in the KO rat is more viscous by 3 months of age (Figure 4D), corresponding to the results seen in MCT rate analysis. Overall, these studies demonstrate that while PCL depth, solid content, and pH are affected at baseline in CF at all ages, they do not contribute to delayed MCT rates until the glands mature, when stimulated PCL collapse and increased mucus viscosity occur in response to secretion of elevated mucus solid content. Thus, mature gland secretion is required to manifest the CF defect primed by airway dehydration and bicarbonate deficiency.

Appropriate mucus properties are dependent on bicarbonate at the surface of the airway. Previous studies, from our laboratory and others (12, 24, 42), have shown that bicarbonate secretion through the CFTR is critical for appropriate mucus transport. To determine whether this was true in the KO rat model and to test the relationship of this finding to mucus hydration, we conducted the same imaging experiments on WT tracheae in the presence and absence of bicarbonate and compared these to KO tracheae. In the absence of bicarbonate, 3-month-old WT tracheae exhibited reduced pH of the airway surface, similar to the KO tracheae (Figure 5A). Interestingly, the lack of bicarbonate transport did not change the solid content percentage (Figure 5B) or PCL depth (Figure 5C) in the 3-month-old WT rat tracheae; this is similar to the effects seen previously in pig tracheae treated with bicarbonate transport inhibitors (24, 28, 42, 43) and is consistent at 1 and 6 months of age (data not shown). In contrast, the effects of absent bicarbonate on MCT rates recapitulated diminished transport in KO tracheae but only after gland maturation; in

Figure 4. Altered mucus transport and viscosity. Mucociliary transport (MCT) rates at baseline conditions (A) and after stimulation with acetylcholine (B). Particle-tracking microphorology (PTM) was used to determine the effective viscosity of mucus at baseline conditions (C), which were 0.06 Hz, and after stimulation with acetylcholine (D). Representative tracks of beads in mucus of each condition at 1 month (E), 3 months (F), and 6 months (G) of age. Scale bar: 1 μm. *P < 0.05. Data were analyzed by 2-way ANOVA. Data in A–D are from 6 animals/group. Data in F and G are representative of 3 independent studies.
bicarbonate-depleted tracheae, MCT rates were reduced at 6 months under baseline conditions (Figure 5D) and also at 3 and 6 months in cholinergically stimulated conditions (Figure 5E). Elevated effective viscosity of the airway surface mucus was also dependent on bicarbonate concentrations; 3-month-old WT tracheae, which were incubated on bicarbonate-free media and were cholinergically stimulated, had more viscous mucus on the airway surface (Figure 5F).

A comparison of MCT rates with corresponding effective viscosity values showed a correlation between the two characteristics ($r^2 = 0.468$, $P < 0.05$, Figure 5G). There was a sharp decrease in MCT rates when viscosity values reached approximately 50 cP. This suggests that there is a threshold of mucus viscosity beyond which the airway is no longer able to transport effectively. In comparison, the correlation between MCT rates and the solids percentage was weak (Figure 5H) as was the correlation between the solids percentage and viscosity (Figure 5I), indicating that solid content does not predict MCT rate or mucus viscosity in the CF airway in the absence of considering the maturity and quality of gland secretions. This comparison also showed that blockade of bicarbonate transport could replicate the effects on mucus viscosity and delayed transport seen in the KO rat tracheae, whereas maturation of the glands was needed to invoke PCL collapse. This indicates that bicarbonate-deficient

Figure 5. Hyperacidic airway surface liquid and altered mucus are bicarbonate dependent. pH of 3-month-old WT tracheae in normal conditions and in bicarbonate-free conditions compared with KO tracheae, was measured using a solid content pH probe (A). Solid content of the airway surface mucus in 3-month-old WT trachea under normal and bicarbonate-free conditions was also compared with KO tracheae (B). Periciliary layer (PCL) depths of WT tracheae in normal conditions and in bicarbonate-free conditions, compared with KO tracheae, obtained from μOCT imaging (C). μOCT imaging also yields mucociliary transport (MCT) rates at baseline (D) and after cholinergic stimulation (E). Particle-tracking microrheology (PTM) yields effective viscosity of WT tracheae in the presence and absence of bicarbonate, compared with KO (F). MCT rates compared with effective viscosity indicate a strong relationship between viscosity of the mucus and the rate at which it is transported across the trachea (G). However, MCT rates compared with the solid content percentage (1-month KO conditions marked by $\tau$) (H) and the solid content percentage compared with viscosity (I) do not correlate in a statistically significant manner. Data are shown as mean ± SEM. *$P < 0.05$, **$P < 0.01$, ***$P < 0.001$, ****$P < 0.0001$. Data were analyzed by (A–C) 1-way ANOVA and (D–F) 2-way ANOVA. Data in A depict 3 regions of interest (ROIs) from 4 animals/group. Data in B are from 3 animals/group. Data in C are 4 ROIs from 6 animals/group. Data in D–I are from 6 animals/group.
mucus exhibits abnormal properties independent of the deleterious effects of airway dehydration. It also indicated that the bicarbonate deficiency in the CF airway may be a therapeutic target to restore appropriate mucus transport.

**Addition of bicarbonate restores normal mucus transport in the CFTR KO rat.** Because bicarbonate concentrations were important to the maintenance of appropriate viscosity and mucus transport, we hypothesized that replacement of bicarbonate on the airway surface would restore aberrant mucus transport in the KO rat tracheae, a concept proposed for CF therapy but not yet tested in vivo (44). These experiments also provided an opportunity to test an intervention independent of airway hydration to define the importance of bicarbonate balance on MCT rates. In tracheae from 6-month-old WT and KO rats, we conducted μOCT imaging at baseline and following addition of sodium bicarbonate at concentrations of 23 mM, 69 mM, 92 mM, and 115 mM to the apical surface of tracheae ex vivo or sodium chloride at 7% (hypertonic saline) as a control for added volume and to compare with a widely used CF therapy (45–47). MCT rates were measured at 5, 20, and 35 minutes following addition of each solution. Compared with baseline, MCT rates in KO rat tracheae increased with each additional concentration of sodium bicarbonate, while MCT rates were unchanged from baseline in WT rat tracheae (Figure 6A). Ciliary beat frequency did not significantly change from baseline in either WT or KO rat tracheae (Figure 6B), indicating that ciliary dynamics were not principally responsible for the increased transport. Hypertonic saline addition to the apical surface also resulted in a significant increase in MCT rate from baseline and was preferentially effective in KO tracheae, but efficacy was less than that when 115 mM bicarbonate was applied (2.53 ± 0.5 mm/min KO plus bicarbonate, 0.92 ± 0.1 mm/min KO plus hypertonic saline, vs. 0.09 ± 0.04 mm/min KO baseline, \( P < 0.001 \) bicarbonate vs. baseline, Figure 6D). Despite substantial efficacy of sodium bicarbonate on MCT rates in KO tracheae, the improvement was transient; the peak effect occurred at 20 minutes after addition and returned to baseline by 35 minutes after addition (Figure 6C). This finding is consistent with the short half-life of bicarbonate at the surface of the airway, even at concentrations much higher than physiologic, likely due to ubiquitous expression of carbonic anhydrase (48, 49).
The effect of bicarbonate addition on MCT rate could be due to altered bicarbonate concentration, alkalinization of the airway pH, or a combination of the two, since the solutions of sodium bicarbonate used here were not pH adjusted. To distinguish these mechanisms, we tested solutions of 115 mM sodium bicarbonate, 50 mM Tris, and 10 mM HEPES and buffered these solutions to pH 6.5, 7.0, 7.5, 8.0, and 8.5. As before, solutions were added to the apical surface of 6-month-old WT and KO rat tracheae, and MCT rates 20 minutes after addition were compared with baseline MCT rates. We found that sodium bicarbonate added at any pH increased MCT rates significantly above baseline in KO tracheae. By contrast, addition of Tris, which has the ability to donate or accept hydrogen ions as necessary, only increased MCT rates when buffered to pH 7.5, which approximated the normal WT pH (approximately 7.2). In contrast, addition of HEPES solution, which, as a zwitterion, has less ability to donate or accept hydrogen ions and therefore is less interactive with its environment, did not alter MCT rates in the KO tracheae from baseline at any pH tested (Figure 7A). These same solutions added to the apical surface of 6-month-old WT tracheae had minimal and nonsignificant effects on MCT rates (Figure 7B). In comparing absolute values of mucus transport in KO tracheae at baseline and 20 minutes after each solution at pH 7.5, both sodium bicarbonate and Tris solutions restored mucus transport in the KO rat tracheae, exceeding MCT rates beyond that observed in the WT tracheae at baseline (3.35 ± 0.9 mm/min KO plus bicarbonate, 3.06 ± 0.9 mm/min KO plus Tris, compared with 0.095 ± 0.02 mm/min KO baseline, P < 0.001, and 0.69 ± 0.2 mm/min WT baseline, Figure 7C). These data indicate that, although the abnormal MCT becomes increasingly severe in the maturing KO rat airway, the defect may be amenable to mucus-targeting therapeutics that restore pH-dependent counterion equilibrium.

**Discussion**

Here for the first time to our knowledge, using complimentary methods and the ability to monitor CF rats over time, we have established a relationship between the development of submucosal glands and the onset of abnormal mucus transport and viscosity in CF. We also demonstrate that both airway dehydration and bicarbonate deficiency independently contribute to the CF mucus defect in this model. Further, we have demonstrated that aberrant mucus transport can be corrected but only by ionic exchange to normalize mucus viscosity. This findings are significant as they lend insight into fundamental aspects of mucus maturation and point to therapeutic interventions to restore mucus transport that may be independent of specific CFTR mutation and, thus, could represent a viable target in all patients with CF.
Detailed investigations in the present study demonstrate that age-dependent changes in submucosal glands may be critical for the KO rat to develop mucus abnormalities. The mechanisms underlying the developmental delay of submucosal glands are unclear but may be related to defective CFTR secretion that manifests over time. The eventual emergence of glandular hyperplasia in the trachea is relevant to human disease, as submucosal glands are thought to be important to airway disease development, as dense mucus secretions and dilated gland ducts were a recognized feature of the CF airways, even early in disease progression (32, 50, 51). The fluid content of the gland secretions is heavily dependent on anion transport and is therefore highly susceptible to dysfunctional CFTR (31). The correlation of glandular hypertrophy with the appearance of the mucus defect in the rat is additional evidence to suggest that the properties of submucosal gland mucus are very important to the CF defect.

While the development of a mucus defect is an important feature of the model, the younger CFTR KO rat offers an opportunity to investigate the relative contributions of the other components of the mucosal surface (before gland maturation). Even in the absence of functional gland secretion, the 1-month-old KO rat exhibits a partial phenotype, with a dehydrated, hyperacidic airway and elevated solid content of the airway mucus; however, mucus transport and viscosity remain normal despite these features. These data suggest that both PCL depletion and hyperacidic pH may be necessary but are not sufficient to cause the CF mucus defect in the absence of mature gland secretions. Mucus secretion, likely principally through glands, is also needed to induce disease. These data may help our understanding of the progression of human CF airway disease, in which airway disease, mucus obstruction, and lung function worsen as the patient ages, as well as our understanding of progression after certain triggers, such as viral or bacterial infection or particle inhalation, when mucus secretion is enhanced (52–55).

Inadequate mucus hydration, reflected by measurement of both PCL depth and the percentage of solids, and abnormal mucus rheology have been suggested as inherent causes of the CF mucus defect (6, 7, 11, 15, 56). These two characteristics interact heavily with one another and are difficult to parse; however, the data presented here suggest that they are not completely dependent upon one another. This is particularly evident in young KO rats, when cholinergic stimulation normalized PCL depth and the percentage of solids, likely through aqueous fluid secretion with few glandular solids. Likewise, the hyperacidic pH at the young age did not confer abnormal mucus transport. Secretions at this young age are hyperacidic and osmotically depleted but not hyperviscous, providing evidence that not all solids are created equal and that they may not all be pH sensitive. This is reflected in the aberrant mucus properties identified by combined evaluation of percentage of solid content and the particle-tracking microrheology methods. In 6-month-old rats, the classical CF phenotype emerges; the PCL collapse under the presence of thicker, glandular secretions is correlated to an emerging mucus transport defect and suggests that a mature mucus load imparts the osmotic pressure imbalance seen in many different models of CF (7, 15). The abundance of mucins secreted from the glands and the role of these to clear the airways emphasize the importance of glandular secretions and may explain why abnormalities in these secretions have such a dramatic effect by 6 months of age (57, 58). This effect can also be recapitulated at an intermediate age (3 months) when gland secretion is stimulated. Taken together, these results strongly suggest that the effects of dehydration and pH are necessary but require the effects of functional airway submucosal glands to cause the complete mucus defect. This may be why CF piglets and neonates are often normal early in life but progressive as infectious insults mount, increasing the stimulus for mucus secretion (59, 60).

Indeed, these mechanisms appear to be distinct when searching for a means of correction of the mucus defect. Merely adding volume to the KO rat trachea, in the case of hypertonic saline, did result in an increase in mucus transport rates, as expected from previous studies using rehydration as a therapeutic technique (61); however the addition of bicarbonate increased transport rates to a much higher degree. By contrast, adjustment of pH only increased transport under specific conditions. Addition of the zwitterionic compound HEPES did not alter mucus transport at all, even at the physiologic pH of 7.5, perhaps because it cannot donate a counterion. In contrast, addition of the ionic Tris buffer at a normal pH, or bicarbonate addition at any pH within the physiologic spectrum, increased MCT rates of CF mucus. These results provide guidance, as mucus normalization strategies are considered for CF therapy, and point out that ionic exchange may be a prerequisite (62). This mechanism may also be contributing to calcium chelation of bicarbonate, an effect that has been demonstrated in other model systems (63). It has been previously shown that both dehydrated mucus and decreased pH independently cause increased viscosity in the CF condition (36, 64), likely without altering other
biophysical factors of the mucus, such as reduction/oxidation states (65); therefore, we hypothesize that the various means of restoration of normality tested here all will result in a reduction of mucus viscosity to more transportable range.

The data presented in this paper offer a model for study of CF airway disease progression, providing a phenotype (before gland development) that may mimic early CF lung disease as well as a state of mucus abnormality (after gland secretion matures) that may more accurately replicate later stages of CF lung disease. Further, data gleaned from these studies indicate that the mechanisms that underlie aberrant mucus movement across the epithelial surface may not fit neatly into one hypothesis but rather involve several factors. This provides a tool to assess what changes as the airway glands mature, allowing the full CF defect to emerge.

**Methods**

**CFTR KO rat model.** All experiments used SD-CFTR<sup>tm1sage</sup> rats (Horizon Discovery), either KO rats or their littermate controls, CFTR<sup>+++</sup> (termed WT) rats. This rat strain was bred and genotyped as previously described (26). In brief, heterozygote (CFTR<sup>+-</sup>) males and females were paired to generate WT and KO pups, and litters remained with lactating dams until 21 days of age. WT and KO rats were maintained on a standard rodent diet, with supplemental DietGel 76A (ClearH20) and 50% Go-LYTLEY (Braintree Laboratories Inc.) added to the water from weaning as a means to reduce mortality from gastrointestinal obstruction. CFTR KO pups have a 50% survival rate under these nutritional conditions but remain up to 30% smaller than their WT littermates by weight. WT and KO rats were assayed at 1, 3, and 6 months of ages, and groups were split evenly between males and females.

**Histology/morphometry.** Tracheae and lungs were immersion fixed in 4% buffered formalin and embedded in paraffin blocks for sectioning. Sections were stained with Alcian blue–periodic acid–Schiff. Slides were imaged; submucosal gland area, intracellular mucus area, goblet cell number, goblet cell area, and intra-airway mucus area were measured morphometrically, as previously described (26).

**μOCT image acquisition and analysis.** Measurements of functional microanatomic parameters in ex vivo tissue were performed using μOCT, a high-speed, high-resolution microscopic reflectance imaging modality. Tracheae were excised and immediately placed on gauze soaked in F12 media, such that the apical surface of the trachea remained media free, and incubated under physiologic conditions (37°C, 5% CO<sub>2</sub>, 100% humidity; using live-cell imaging incubation systems, ref. 24; Carl Zeiss). Tracheae were allowed to equilibrate for 1 hour before imaging. Following baseline image acquisition, tracheae were moved onto F12 media-soaked gauze with 100 μM acetylcholine added and incubated again for 30 minutes at physiologic conditions and reimaged. We have previously described μOCT methods for interrogation of airway epithelia and quantitative image analysis, including calibration of distance and height measurements (34). Images were acquired at 5 controlled points along the tracheal length, using the cranial end as a reference. MCT rate was determined using time elapsed and distance traveled of native particulates in the mucus over multiple frames, using the pixel-to-micron conversion factor determined with image calibration. For each trachea, images were acquired at standard distances along the ventral surface, with the optical beam scanned along the longitudinal direction.

**Measurement of the solid content percentage.** Tracheae were prepared as above. Filter paper was cut to a size that would fit inside an open tracheae. Papers were weighed, placed on each trachea for 15 seconds, and immediately reweighed to determine mucus collection. Filters were left overnight in a dry oven for liquid evaporation and weighed again for measurement of solid content. All weights were taken using a UMX2 Ultra-microbalance (Mettler Toledo). Filter weights were subtracted, leaving wet and dry weights of the mucus, which were used to calculate the fraction that was solid content.

**Particle-tracking microrheology.** Particle-tracking microrheological techniques were used to measure viscosity of mucus on rat tracheae in situ. Tracheae were treated with 0.1% benzalkonium chloride (Acros Organics) for 1 hour at 37°C to stop ciliary beating. Polystyrene beads were used as previously described (24, 40). Baseline μOCT images were acquired; tracheae were incubated for 30 minutes at 37°C on media with 100 μM acetylcholine; and then stimulated μOCT images were acquired. Images were analyzed using ImageJ (NIH) and the SpotTracker plug-in (66). Resulting particle tracks were analyzed with custom Matlab procedures to compute mean squared displacement (MSD), while subtracting spurious bulk motion common to all tracks. Dynamic viscosity was derived from MSD by application of the generalized Stokes-Einstein relation as described previously (24, 40).
Transport inhibition. In some experiments, 100 μM acetylcholine was added basolaterally to the F12 media for cholinergically stimulated mucus release. For experiments that blocked chloride transport, 100 μM bumetanide was added to F12 media; bicarbonate transport was inhibited with use of F12 media without sodium bicarbonate. Tracheae were incubated under all conditions for 30 minutes before μOCT imaging. Bicarbonate-free experiments were incubated in 0% CO₂.

Apical addition. For MCT rescue experiments, 5 μl of each solution was added to the distal end of the apical surface of the tracheae and allowed to equilibrate before μOCT imaging at 5, 20, and 35 minutes after addition. Solutions added were sodium bicarbonate at concentrations of 23 mM, 69 mM, 93 mM, and 115 mM (non-pH adjusted); 0.9% saline (non-pH adjusted); 7% saline (HTS, non-pH adjusted); 115 mM sodium bicarbonate adjusted to a pH of 6.5, 7.0, 7.5, 8.0, or 8.5; 50 mM Tris-buffered saline, adjusted to a pH of 6.5, 7.0, 7.5, 8.0, or 8.5; and 100 mM HEPES, adjusted to a pH of 6.5, 7.0, 7.5, 8.0, or 8.5.

Statistics. Statistical analysis was performed with GraphPad Prism version 7.0. Inferential statistics (mean, SD, and SEM) were computed using ANOVA or 2-tailed unpaired or paired t test, as appropriate. For multiple comparisons, post-hoc testing was applied only if ANOVA was significant. P values of less than 0.05 were considered significant. Statistics are presented as mean ± SEM, unless indicated otherwise. Except where noted, mean values per trachea explant are reported.

Study approval. All animal use was approved by the University of Alabama at Birmingham IACUC.

Author contributions
SEB, KLT, M VF , EJS, and SMR conceived of the experiments; SEB, CMF, KLT, AMO, and JMD performed research; SEB, CMF, KLT, and KKC analyzed the data; SEB, GJT, M VF, EJS, and SMR contributed reagents, materials, and/or analysis tools; SEB and SMR wrote the manuscript; and GJT, M VF, EJS, and SMR supervised the project.

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