Mechanical ventilation is an essential lifesaving therapy in acute respiratory distress syndrome (ARDS) that may cause ventilator-induced lung injury (VILI) through a positive feedback between altered alveolar mechanics, edema, surfactant inactivation, and injury. Although the biophysical forces that cause VILI are well documented, a knowledge gap remains in the quantitative link between altered parenchymal structure (namely alveolar derecruitment and flooding), pulmonary function, and VILI. This information is essential to developing diagnostic criteria and ventilation strategies to reduce VILI and improve ARDS survival. To address this unmet need, we mechanically ventilated mice to cause VILI. Lung structure was measured at three air inflation pressures using design-based stereology, and the mechanical function of the pulmonary system was measured with the forced oscillation technique. Assessment of the pulmonary surfactant included total surfactant, distribution of phospholipid aggregates, and surface tension lowering activity. VILI-induced changes in the surfactant included reduced surface tension lowering activity in the typically functional fraction of large phospholipid aggregates and a significant increase in the pool of surface-inactive small phospholipid aggregates. The dominant alterations in lung structure at low airway pressures were alveolar collapse and flooding. At higher airway pressures, alveolar collapse was mitigated and the flooded alveoli remained filled with proteinaceous edema. The loss of ventilated alveoli resulted in decreased alveolar gas volume and gas-exchange surface area. These data characterize three alveolar phenotypes in murine VILI: flooded and non-recruitable alveoli, unstable alveoli that derecruit at airway pressures below 5 cmH$_2$O, and alveoli with relatively normal structure and function. The fraction of alveoli with each phenotype is reflected in the proportional changes in pulmonary system elastance at positive end expiratory pressures of 0, 3, and 6 cmH$_2$O.

Keywords: ventilator-induced lung injury, stereology, pulmonary surfactant, lung function, alveolar mechanics
INTRODUCTION

Mechanical ventilation is required for major surgery and resuscitation and in critical emergencies such as acute respiratory distress syndrome (ARDS) (Force et al., 2012), but this lifesaving therapy carries the risk of ventilator-induced lung injury (VILI). The mechanisms of VILI are well documented and include direct tissue injury from overdistension (volutrauma) (Dreyfuss and Saumon, 1998; Hamlington et al., 2018b), allowing the ingress of proteinaceous edema into the distal airspace that is associated with changes in lung mechanics (Hamlington et al., 2016a).

Changes in the lung structure-function relationship during VILI are currently understood through the lens of organ-scale imaging techniques such as CT (Yen et al., 2019). However, the positive feedback mechanism between leak, surfactant function, altered mechanics, and injury occurs at the alveolar and acinar scales (Knudsen et al., 2018). As such, measurements of morphometry are necessary to bridge studies of cellular injury and alveolo-capillary barrier disruption (Hamlington et al., 2018a, b) to organ-scale measurements of structure and function. Understanding how these mechanisms are associated across length scales is critical for developing approaches to mitigating them. Structure-function correlations may also improve clinical practice by defining the mechanistic basis for diagnostic parameters such as the driving pressure (Aoyama et al., 2018; Pereira Romano et al., 2020) and subject-specific computational models (Hamlington et al., 2016b; Mellenthin et al., 2019; Morton et al., 2019) that seek to optimize ventilation based on patient-specific alveolar mechanics.

To address this knowledge gap, we used design-based stereology to measure parenchymal structure over a range of air inflation pressures in a mouse model of VILI. The state of the pulmonary surfactant system was assessed by measuring minimum surface tension and surfactant composition. Organ-scale mechanical function was determined with the forced oscillation technique (FOT) to quantify stiffness, airflow resistance, and tissue damping at different levels of lung inflation. Together, these data define the structure–function relationship in murine VILI.

MATERIALS AND METHODS

Animal Procedures

Eight to ten week old female BALB/c mice (Jackson Laboratories, Bar Harbor, ME, United States) weighing 17.1–22.3 g were studied under Institutional Animal Care and Use Committee (IACUC)-approved protocols. Mice were anesthetized with an intraperitoneal (IP) injection of 100 mg/kg ketamine and 16 mg/kg xylazine, tracheostomized with a 18 ga metal cannula, and ventilated using a flexiVent small animal ventilator (SCIREQ, Montreal, QC, Canada). Respiratory drive was suppressed via 0.8 mg/kg pancuronium bromide administered at the onset of mechanical ventilation. Alternating doses of 50 mg/kg ketamine and 50 mg/kg ketamine with 8 mg/kg xylazine were administered at 30-min intervals with 150 µL IP 5% dextrose lactated Ringer’s solution. Fluid volume from the anesthetic was 10 µL/g. The electrocardiogram was monitored to assure that the rodents maintained a deep plane of anesthesia during paralysis.

Mechanical ventilation experiments to generate tissue for structural analysis were conducted at the University of Vermont and approved by the University of Vermont Institutional Animal Care and Use Committee (IACUC #14-056). Bronchoalveolar lavage fluid for surfactant analysis was collected under identical ventilation conditions by the same investigator at the University of Colorado Denver in experiments approved by the University of Colorado Denver Anschutz Medical Campus, IACUC (#00230).

Ventilation Protocol

All animals were ventilated for a 10-min stabilization period with a delivered tidal volume (Vt) = 10 ml/kg at a respiratory rate (RR) = 200 breaths/min and a positive end expiratory pressure (PEEP) = 3 cmH2O (baseline ventilation). Recruitment maneuvers (RM) consisting of a 3 sec ramp to 30 cmH2O followed by a 3 sec breath hold were applied to re-open regions that may have collapsed during the surgical procedure. Following the stabilization period, lung function was assessed. Testing consisted of an RM and a dynamic pressure-volume (PV) loop that ramped airway pressure (Paw) from 0 to 37.5 cmH2O, held pressure for 3 sec, and ramped pressure back to zero over 3 sec. Derecruitability tests were then performed at PEEP = 6, 3, and 0 cmH2O, denoted by $D_{rec}^{PEEP6}$, $D_{rec}^{PEEP3}$, and $D_{rec}^{PEEP0}$. Each test consisted of an RM and then Vt = 10 ml/kg and RR = 200 breaths/min at the specified PEEP. Nine multifrequency FOT impedance measurements were recorded at 18 s intervals and fit to the constant phase model (Hantos et al., 1992) to determine respiratory system elastance (H), tissue damping (G), and Newtonian resistance (Rn). The total duration of the lung function assessment was 13 min. Lungs from the control group were then harvested for morphometric analysis. In a separate sub-group of animals, bronchoalveolar lavage fluid (BALF) was collected for surfactant analysis.

The VILI group was then subjected to repeated blocks of ventilation consisting of 4 min 45 s of injurious ventilation with a plateau pressure (Pplat) = 37.5 cmH2O, RR = 50 breaths/min, inspiratory:expiratory ratio of 1:2, and PEEP = 0 cmH2O. This Pplat was selected to generate severe VILI in an experimentally tractable timeframe without causing any pneumothoraces. At the end of each block a PV loop was recorded followed by two FOT measurements separated by 10 s of baseline ventilation. These 5-min ventilation blocks were applied until H doubled, at which point the lung function assessment was repeated prior to harvest for either morphometry or surfactant analysis. The mean duration of injurious mechanical ventilation was 87.3 min with a standard deviation of 17.5 min. Including with the 10 min stabilization period and two 13 min lung function assessments, the mean duration of ventilation for the VILI group was 123.3 min.
Surfactant Analysis

Bronchoalveolar lavage fluid was collected by instilling and suctioning back 1 ml warm normal saline three times and then centrifuging at 700 g for 5 min. A total of 100 µl of the cell-free supernatant for each animal was separated, and the remainder of the BALF from three mice in the same treatment group was combined and centrifuged at 40,000 g. The ultracentrifuged supernatant containing the surfactant small aggregate (SA) was stored at −80°C for later analysis. The surfactant pellet, containing the large aggregate (LA) fraction, was washed twice by resuspending the pellet in 1 ml saline and centrifuging at 40,000 g. Finally, the LA pellet was resuspended in 100 µl saline and stored at −80°C for later analysis.

Quantitation of the surfactant recovered from the BAL was completed by measuring lipid phosphorous content in the LA and SA fractions as previously described (Rouser et al., 1970). Total surfactant recovery represents the combined amount of both fractions. Total BAL protein was quantified with a BCA assay (ThermoShandon-Pierce). Surfactant surface lowering activity was assessed by measuring minimal surface tension of the LA fraction ex vivo using a pulsating bubble surfactometer (General Transco) as described (Enhorning and Holm, 1993; Hite et al., 2005). Surfactant was preheated to 37°C at 2 mg (PL from LA)/ml in 150 mM NaCl, 5 mM CaCl₂, 5 mM Tris solution, and the samples were pulsed at 20 pulses/min for 20 min.

Lung Tissue Preparation

Lung structure was analyzed in a separate subgroup of mice that were not subjected to a bronchoalveolar lavage. The lungs were fixed through the vasculature while the air inflation pressure was held at a prescribed level (described below) to maintain surface tension effects and allow comparison to the lung function data (Gil et al., 1979; Bachofen et al., 1982). The fixation process was initiated during baseline ventilation by performing a bilateral thoracotomy and flushing the pulmonary circulation with 5 ml 1% OsO₄ in 0.15 M HEPES buffer for approximately 5 min before immersion fixation for at least 24 h.

Fields of view were selected for analysis using systematic uniform random sampling (Knudsen et al., 2018) where the sampling fraction, Fₘ, was 2.8% at magnification (Fₘ = 2.8%), 1,200 μm grid spacing) by 10% of the images (distributed randomly across groups and subjects) were recorded using an Olympus BX53 with a DP73 camera (Olympus, Waltham, MA, United States) controlled with the NewCast stereology software (Visiopharm, Hørsholm, Denmark). Stereological quantification of all images was performed using a custom MATLAB (Mathworks, Natick, MA, United States) graphical user interface.

A cascade sampling design was used starting with the volume fraction of parenchyma in the whole lung [Vv(par/lung)] that we define as areas contributing to gas exchange and excluding airways, vessels outside the septal walls, and peribronchiolar tissue. This, and all volume fractions, were measured using point counting. The percentage of the tissue section analyzed (the sampling fraction, Sₚ) for Vv(par/lung) was 100% at 5× magnification. The parenchyma was then subdivided into volume fractions of alveolar airspace [Vv(alvair/par)], alveolar duct airspace [Vv(ductair/par)], and non-air material [Vv(non-air/par)] that include tissue and airspace edema fluid. This assessment was performed at 20× magnification and Sₚ = 11%.

Fields of view were selected for analysis using systematic uniform random sampling whereby the morphology software selected a random starting location and images were then automatically sampled on a 1,200 × 1,200 μm grid over the entire tissue section. This approach blinks the investigator to the selection of the images used for analysis, thus removing the potential for bias. The non-air material in the parenchyma [Vv(non-air/par)] was then analyzed at 40× magnification (Sₚ = 2.8%, 1,200 μm grid spacing) to determine the volume fractions of sepal tissue of patent alveoli [Vv(sep,air/par)], total sepal tissue [Vv(sep,total/par)], and airspace edema [Vv(edema/par)]. The volume of each compartment (demarcated with V in place of Vv) was determined by multiplying the volume fractions by the volume of the reference space. The surface area per volume available for gas exchange [Sv(alvair/par)] was estimated at 40× magnification (Sₚ = 2.8%, 1,200 μm grid spacing) by counting line intersections with aerated septal tissue that was not covered with alveolar edema. The gas exchange surface area
Pulmonary system elastance (H) was determined by multiplying Sv(alvair/par) by the parenchymal volume [V(par)] and the mean septal thickness defined as τ(sep) = 2 V(sep,air)/S(alvair). The physical dissector method (Ochs et al., 2004) was used to determine the number of patent [N(alv)] and flooded [N(flood)] alveoli that we define as alveoli with visible architecture and alveolar edema fully blocking the alveolar mouth using 40× magnification (Sf = 6.25%, 800 μm grid spacing). The total number of alveoli is then N(alv,total) = N(alv) + N(flood), the number-weighted average alveolar volume \( V_n(alv) = V(alvair)/N(alv) \), and the alveolar surface area to volume ratio \( S/V(alv) = S(alvair)/V(alvair) \).

### Statistical Analysis

Data were curated in MATLAB and statistical analysis was performed in R (The R Foundation). A Shapiro–Wilks test was used to assess normality. If the data were not significantly different from the normal distribution \( (p > 0.05) \) then an ANOVA was performed followed by pairwise comparisons using the lsmeans package (Lenth, 2016) and Tukey’s adjustment for multiple comparisons. Data that failed the normality test was analyzed with a Kruskal–Wallis test followed by a rank-sum Conover–Iman test (PMCMR package) with the Holm correction for multiple comparisons.

The time series of lung function parameters H, G, and Rn measured during the derecruitability tests were analyzed using a linear mixed effects model (LME) (NLME package in R) on the natural logarithm of the time and parameter values. This is equivalent to fitting the equation \( y = \alpha t^\beta \) to the time series of measurements as we have previously described (Smith et al., 2017). Fixed effects were derecruitability test PEEP \( (D_{PEEP}) \), the time within the derecruitability test \( (t) \), and whether the measurement was recorded before or after VILI \( (Injury) \). The intercept for each animal was included as a random effect and the Akaike information criterion (AIC) was used to determine the most appropriate model. Temporal autocorrelation was accounted for with a first-order autocorrelation structure for the continuous time covariate grouped by animal, derecruitability test PEEP, and injury status (Pinheiro and Bates, 2000).

### RESULTS

Pulmonary system elastance following an RM in the uninjured lungs (Figure 1, first row) was significantly different between the derecruitability tests at different PEEP levels, while the rate of change of H over time \( (the slope) \) remained consistent between PEEPs (Table 1). VILI (Figure 1, second row) increased the H intercept at all PEEPs with a proportionally greater increase occurring in \( D_{PEEP} \) where the elastance at the start of the derecruitability test was approximately doubled. The rate of elastance increase over time was also elevated in \( D_{PEEP} \) and, to a lesser extent, in \( D_{PEEP} \). Tissue damping \( (G) \) increased monotonically with PEEP prior to injury (Figure 2, first row) and was independent of \( t \) (Table 2). Following VILI, G demonstrated a marked increase in \( D_{PEEP} \) that was similar in magnitude to the change in H. Central airway resistance \( (Rn) \) tended to decrease with PEEP (Figure 3) in both healthy and injured lung. Following VILI, Rn was slightly increased in \( D_{PEEP} \) and decreased in \( D_{PEEP} \). The temporal dependence of Rn was not affected by VILI (Table 2).

The changes in lung function are associated with structural alterations in the perfusion-fixed lung tissue. Alveoli in the Control lungs (Figure 4, first row) remained patent from Paw = 10 cmH\(_2\)O (right column) down to airway pressures of 2 cmH\(_2\)O (left column). Note that these airway pressures are applied in open-chested mice, and the contributions of the chest wall are discussed below. By contrast, the VILI lungs exhibited diffuse alveolar edema (second row, asterisks) and retraction of the alveolar septa (second row, arrows) at Paw = 5 and 10 cmH\(_2\)O. At Paw = 2 cmH\(_2\)O, the injured parenchyma tended to consolidate into patches of flooded and collapsed alveoli with air remaining in the ductal space. These injured regions were interspersed between areas of parenchyma that appeared normal.

Quantitative structural changes with VILI included a significant increase in V(alvair) (Figure 5A) in conjunction with reduced N(alv) (Figure 6B). The average alveolar volume \( [V_n(alv)] \) is defined as the ratio of these two quantities and indicates a modest stiffening of patent alveoli with injury (Figure 6E). By contrast, V(ductair) did not demonstrate significant alterations in VILI (Figure 5B), leading to a fractional shift of ventilation volume from the alveoli to the alveolar ducts with VILI (Figure 5C) that was significant at all airway pressures. The surface area available for gas exchange (Figure 5D) was significantly reduced across all airway pressures in VILI. This reduction is primarily due to a loss of ventilated alveoli and not to reduced surface area to volume ratio in the remaining patent units as indicated by increased S/V(alv) when comparing control and VILI at Paw = 2 cmH\(_2\)O (Figure 6F, lines). For comparison, Figure 6F shows the hypothetical surface area to volume ratios for perfectly spherical alveoli (symbols), and these values are less than those measured in vivo despite the reduction in area caused by the alveolar mouth. The increase in S/V(alv) at low Paw in VILI suggests that the alveolar geometry becomes more distorted in the injured lung. Injurious ventilation did not change V(sep, total) (Figure 5E) or t(sep), and although septal thickness tended to increase with decreasing Paw there were no significant alterations (Figure 6A). Little to no airspace edema was observed in the control animals (Figure 5F), and V(edema) was constant across inflation pressures in VILI.

The number of ventilated alveoli \( [N(alv)] \), Figure 6B) remained constant across inflation pressures in the Control animals, indicating an absence of derecruitment even at Paw = 2 cmH\(_2\)O. The VILI mice demonstrated a constant N(alv) at Paw = 5 and 10 cmH\(_2\)O that was significantly less than in the Controls. Reducing the inflation pressure to 2 cmH\(_2\)O in the VILI animals led to a significant reduction in aerated alveoli. Flooded alveoli \( [N(flood)] \), Figure 6C) were defined as having a liquid meniscus fully spanning the alveolar mouth. N(flood) remained constant across inflation pressures in the VILI group, and no flooded alveoli were observed across the 15 control lungs analyzed (Table 3). Figure 6D shows that the total number of flooded and aerated alveoli \( [N(alv,total)] \) is consistent across all
The numeric values for slope indicate the change in Δ from the left-hand to the right-hand derecruitability test listed under “Pairwise Comparison.” The numeric values provided for the intercepts indicate the fold change in the raw (not log transformed) intercept α. Gray cells indicate p < 0.05. Fifteen mice are in each group.

The changes in lung structure and function are accompanied by alterations in the pulmonary surfactant system. BALF total phospholipid (PL) content (Figure 8A) significantly increased correlated with the number of patent alveoli ($R^2 = 0.78$), and the form of the regression was selected based on our prior computational simulations where elastance increases with the inverse of the lung open fraction (Smith et al., 2013, 2015; Hamlington et al., 2016b; Mellenthin et al., 2019).
TABLE 2 | Results from regression analysis predicting log-transformed H, G, or Rn from derecruitability test PEEP (DEEP0) with factors DEEP0rec, DEEP3rec, and DEEP6rec; the log-transformed continuous time within the derecruitability test (t); and an Injury factor describing if the measurements were recorded at the start of the experiment (pre-ventilation) or after ventilation to a doubling of pulmonary system elastance (VILI).

| Predictor | Estimate (SE) | p-value | Estimate (SE) | p-value | Estimate (SE) | p-value |
|-----------|---------------|---------|---------------|---------|---------------|---------|
| Intercept | 3.083 (0.034) | <0.0001 | 0.500 (0.062) | <0.0001 | -1.103 (0.057) | <0.0001 |
| t         | 0.038 (0.006) | <0.0001 | -0.002 (0.015) | 0.8857 | 0.035 (0.009) | 0.0002 |
| DEEP0rec  | -0.386 (0.036) | <0.0001 | 0.228 (0.075) | 0.0024 | -0.499 (0.009) | <0.0001 |
| DEEP3rec  | -0.229 (0.036) | <0.0001 | 0.580 (0.075) | <0.0001 | -0.657 (0.047) | <0.0001 |
| DEEP6rec  | -0.229 (0.036) | <0.0001 | 0.778 (0.075) | <0.0001 | 0.146 (0.047) | 0.0021 |
| t × DEEP0rec | -0.012 (0.008) | 0.1529 | 0.002 (0.021) | 0.8910 | -0.016 (0.013) | 0.2292 |
| t × DEEP3rec | -0.014 (0.008) | 0.1088 | 0.001 (0.021) | 0.9627 | -0.032 (0.013) | 0.0138 |
| t × DEEP6rec | 0.060 (0.008) | <0.0001 | 0.115 (0.021) | <0.0001 | 0.008 (0.013) | 0.5550 |
| t × VILI   | -0.309 (0.051) | <0.0001 | -0.710 (0.106) | <0.0001 | -0.176 (0.067) | 0.0089 |
| t × DEEP0rec × VILI | -0.031 (0.012) | 0.0100 | -0.093 (0.030) | 0.0015 | 0.002 (0.019) | 0.9034 |
| t × DEEP3rec × VILI | -0.051 (0.012) | <0.0001 | -0.097 (0.030) | 0.0009 | -0.009 (0.019) | 0.6143 |

Fifteen mice are in each group.

FIGURE 2 | Tissue damping (G) measured during derecruitability tests at PEEP = 0 (A,D), 3 (B,E), and 6 cmH2O (C,F) (columns) before (A–C) and after injurious mechanical ventilation (D–F). The LME model fit is shown with a black line, the 95% confidence interval is shown with a blue band, and the 95% prediction interval is shown with a red band. Symbols depict raw data for each animal and are consistent between Figures 1–3. Fifteen mice are in each group.

following high tidal volume ventilation due primarily to an increase in SA, which are less surface active. The increase in SA fraction also resulted in a striking and significant decrease in the LA/SA ratio (Figure 8B), a frequent marker of surfactant degradation, turnover, and/or injury. The fraction of the most surface-active LA was unchanged after VILI, but the functional quality of the LA pellet was significantly disrupted as demonstrated by a marked increase in minimum surface tension.
(Figure 8D), which translates in vivo to reduced surface tension lowering activity and reduced overall surfactant function. The airspace of the injured lung contained eightfold more protein (Figure 8C), which likely represents a combination of proteins released by the lung parenchyma and serum protein extravasation across a disrupted alveolocapillary barrier.

**DISCUSSION**

The macroscale pressures and flows applied at the trachea during mechanical ventilation result in microscale parenchymal injury (Dreyfuss and Saumon, 1998). The resulting ingress of protein-rich edema into the parenchymal airspace causes changes in alveolar dynamics at the microscale that are then reflected in macroscale alterations in lung function (Hamlington et al., 2018b). Macro- to micro-scale interactions in VILI thus take place in both directions, which we investigated by correlating VILI-induced changes in mouse lung microstructure to changes in lung function. Understanding the link between structure and function at both micro and macro levels of scale is crucial for developing optimized mechanical ventilation strategies that interrupt the vicious cycle of surfactant degradation and inactivation (Agassandian and Mallampalli, 2013), mechanical injury, inflammation, and altered lung function that drives VILI pathogenesis.

The most striking alterations in VILI-related organ-scale function are seen at low lung volumes (Figure 1, column 1), where we observe an approximately twofold increase in the first value of elastance measured at PEEP = 0 cmH\textsubscript{2}O immediately following an RM in VILI compared to controls (Table 1, intercept). This is attributable to the loss of roughly half the ventilated alveoli \([N(alv), \text{Figure 6B}]\), due both to derecruitment that is reversible with an RM and the flooding of alveoli \([N(flood), \text{Figure 6C}]\) that are not recruitable. \(H\) then continues to increase over the subsequent three minutes of \(D\text{PEEP}\) at a rate that is greater in the injured lungs (Table 1, slope). Although we cannot assess time-dependent alveolar collapse in fixed tissue, our previous studies indicate that this increased rate of lung stiffening is a consequence of accelerated alveolar derecruitment (Smith et al., 2015) that we would ascribe, in the current study, to elevated minimum surface tension (Figure 8D).

The PEEP = 3 and 6 cmH\textsubscript{2}O derecruitability tests also show progressive increases in \(H\), although of substantially smaller magnitudes than at PEEP = 0 cmH\textsubscript{2}O (Table 1), no doubt because airway pressures remained above the derecruitment pressures of most of the unstable alveoli at the higher PEEP levels. This is borne out by our stereological analysis, which shows a decrease...
**FIGURE 4** | Representative micrographs for control (first row) and VILI (second row) mice that were perfusion fixed at an airway pressure $P_{aw} = 2$ cmH$_2$O (first column), $P_{aw} = 5$ cmH$_2$O (second column), and $P_{aw} = 10$ cmH$_2$O (third column). Asterisk indicates alveolar edema and arrows denote retracted alveolar septa.

**FIGURE 5** | Volume of air in the alveoli $V_{(alv)_{air}}$ (A), volume of air in the alveolar ducts $V_{(duct)_{air}}$ (B), volume fraction of parenchymal air in the alveolar ducts $V_{f(duct)}$ (C), gas-exchanging surface area $S_{(alv)_{air}}$ (D), total septal tissue volume $V_{(sep,tot)}$ (E), and alveolar edema volume $V_{(edema)}$ (F) for the control (black lines) and VILI (gray lines) mice fixed at airway pressures $P_{aw} = 2, 5,$ and 10 cmH$_2$O. Significant differences from $P_{aw} = 2$ cmH$_2$O (#), $P_{aw} = 5$ cmH$_2$O (%), $P_{aw} = 10$ cmH$_2$O (&) in the same treatment group. Significant difference at the same airway pressure between treatment groups is shown with asterisks. $n = 5$ for each point and the error bars show standard error.
in $N(\text{alv})$ at the higher PEEP levels that is commensurate with the corresponding rates of increase in $H$. However, the total number of alveoli, $N(\text{alv, total})$, which includes both patent and flooded units, was unchanged in the VILI animals compared to controls at $P_{aw} = 5$ and 10 cmH$_2$O. This indicates that the increase in elasstance in $D^{\text{PEEP}}$ and $D^{\text{PEEP}}$ is primarily due to the accumulation of flooded, non-recruitable alveoli rather than recruitment-resistant atelectasis or small airway closure. Figure 7 shows that the $H$ values measured at the end of the derecruitability tests in all animals are inversely related to the fraction of patent alveoli (the “open fraction”), as we have previously postulated in our modeling studies of ventilator and acute lung injury in rodents (Smith et al., 2013, 2015; Hamlington et al., 2016b). This type of relationship was also observed during the acute phase of bleomycin injury in rats (Lutz et al., 2015; Knudsen et al., 2018) and lends strong support to the notion that the dynamics of recruitment and derecruitment in the injured lung may be inferred directly from observed changes in $H$.

As expected, the loss of ventilated alveoli with VILI was accompanied by corresponding losses in $V(\text{alv})$ at all pressures (Figure 5A). Less expected was the reduction in $V_{\text{n,alv}}$ (Figure 6E), because we previously observed mean alveolar volume to increase during the first 3 days of bleomycin injury in rats (Knudsen et al., 2018). Bleomycin-treated rats experience similar changes in minimum surface tension (Lutz et al., 2015) to those we found in the present study, so the differences in alveolar volume behavior cannot be attributed to differences in surfactant function. On the other hand, there may have been differences in the way that alveoli became derecruited in the two models. Finite element simulations indicate that the degree of alveolar enlargement caused by tethering forces from adjacent derecruited regions of the lung is greater when the adjacent alveoli are collapsed as opposed to being merely flooded (Albert et al., 2019). The later stages of bleomycin injury are characterized by widespread alveolar collapse that is resistant to reopening even at high airway pressures (Knudsen et al., 2018), which may explain why the mean volume of open alveoli was increased in this model. In contrast, in the VILI model of the present study the collapsed alveoli could be reopened by an RM, and the alveoli remained patent down to $P_{aw} = 5$ cmH$_2$O, which would have reduced the tethering forces. Furthermore, at $P_{aw} = 2$ cmH$_2$O the VILI lungs were characterized by extensive areas of both flooded and collapsed alveoli interspersed with regions of normal-appearing alveoli, so that tethering-induced alveolar volume increases would be confined to the boundaries between the collapsed and open regions.

The surface area available for gas exchange (Figure 5D) followed the same trend as $V(\text{alv})$, increasing with $P_{aw}$ and decreasing with VILI. However, the mean surface area-to-volume ratio of an individual alveolus (Figure 6F) showed a striking increase with VILI at $P_{aw} = 2$ cmH$_2$O, indicating that the
TABLE 3 | Summarized stereological data from control and VILI lungs.

| Paw cmH₂O | N(alv) | N(flood) | V(alv) | V(edema) | V(alv,air) | V(alv) (air) | N(alv,air) | N(alv,air) | N(alv,air) | N(alv,air) | N(alv,total) |
|-----------|--------|---------|--------|----------|-----------|-------------|------------|------------|------------|------------|--------------|
| Control   | 5      | 0.116   | 0.101  | 0.007    | 0.019     | 0.018       | 0.017      | 0.017      | 0.017      | 0.017      | 0.007        |
| 2         | 5      | 0.116   | 0.101  | 0.007    | 0.019     | 0.018       | 0.017      | 0.017      | 0.017      | 0.017      | 0.007        |
| 5         | 5      | 0.116   | 0.101  | 0.007    | 0.019     | 0.018       | 0.017      | 0.017      | 0.017      | 0.017      | 0.007        |
| 10        | 5      | 0.116   | 0.101  | 0.007    | 0.019     | 0.018       | 0.017      | 0.017      | 0.017      | 0.017      | 0.007        |
| VILI      | 5      | 0.116   | 0.101  | 0.007    | 0.019     | 0.018       | 0.017      | 0.017      | 0.017      | 0.017      | 0.007        |

Lungs were perfusion fixed at airway pressures Paw = 2, 5, and 10 cmH₂O in the same treatment group. Significant difference from Paw = 2 cmH₂O in the controls. Zone D comprises of damaged alveoli that cannot be recruited, represented by the decrease in N(alv) between Paw = 5 and 2 cmH₂O in VILI (Figure 6D). This represents an VILI-induced upward shift of the distribution of alveolar derecruitment pressures (i.e., alveolar instability) since there is no change in N(alv) between Paw = 5 and 2 cmH₂O in the controls. Zone H comprises the remaining open alveoli (i.e., the “baby lung”) that remain aerated at low Paw. Our microscale analysis shows that the mechanics of these open alveoli are not entirely normal since Vₐ(alv) and S/V(alv) are altered in VILI, but these alterations are secondary contributors to organ-scale mechanics when compared to the loss of ventilated alveoli. These three alveolar phenotypes have different dynamic behaviors that are reflected in the time- and pressure-dependence of H. It is important to note that while these three alveolar phenotypes are generally stable there may be variability between respiratory cycles. In vivo confocal microscopy shows that flooded alveoli are typically stable but may occasionally expel their edematous contents and become re-aerated (Wu et al., 2017). Likewise, nearby alveoli may alternately collapse on subsequent respiratory cycles (Broche et al., 2017) so that on subsequent breaths a single alveolus may either be collapsed or open.

The correlation between alveolar recruitment and elastance (Figure 7) may provide a mechanistic basis for identifying optimally lung-protective ventilation strategies. Figure 6 shows alveolar recruitment occurring between pressures of 2 and 5 cmH₂O and, as such, H decreases as PEEP is increased from 0 to 3 cmH₂O (Figure 1). By contrast, further increasing PEEP to 6 cmH₂O causes a modest increase in elastance (and thus driving pressure) that is indicative of parenchymal overdistension. Taken together, these data suggest that the optimal PEEP based on lung elastance (or driving pressure) for these VILI mice lies between 3 and 6 cmH₂O to recruit the recruitable alveoli (Zone R) without causing tissue overdistension by applying excessive PEEP in a futile attempt to recruit the Zone D alveoli that, in our VILI model, are flooded with edema.

Tissue damping is challenging to interpret through the lens of fixed tissue because it describes the viscous losses in the parenchyma that are, by definition, dynamic processes. The fractional changes in G during the derecruitability tests were, for the most part, rather similar to the corresponding fractional changes in H (Table 1), which is most readily interpreted as being...
due simply to derecruitment of lung units, although it could also reflect an intrinsic coupling of the elastic and dissipative properties of lung parenchyma (Kaczkka and Smallwood, 2012). Nevertheless, the largest alteration in G caused by VILI was an approximately twofold increase during $D_{\text{rec}}^{\text{PEEP}}$ (Figure 2D) that was slightly larger than the corresponding relative increase in H (Figure 1D). This may have been due to an increase in some form of dissipative process in the lung, such as intratidal recruitment (Kaczka et al., 2005, 2011), movement of fluid within and between partially flooded alveoli, or the folding and unfolding of septal pleats as suggested by an increased S/V(alv). In contrast, at PEEP = 3 and 6 cmH₂O the VILI-induced changes in G were relatively less than those in H. This contrasts with findings in open-chest but otherwise healthy mice in which tissue hysteresivity = G/H remained constant with lung volume (Sly et al., 2003). Theoretical analysis, however, shows that reductions in hysteresivity can arise if ongoing recruitment and derecruitment of lung units occurs while lung impedance is being measured by the FOT (Bates and Allen, 2006), as might easily be the case in severely injured lungs.

The decrease in airway resistance with PEEP (Figure 3) is readily attributable to the increase in airway caliber caused by airway-parenchymal tethering forces. This mechanism may also explain the modest VILI-induced decrease in $R_n$ during $D_{\text{rec}}^{\text{PEEP}}$, since flooded alveoli near the airways would be unable to expand normally during lung inflation and thus would cause tethering-forces on the airway to be correspondingly increased. Similarly, reduced or compressive tethering forces in atelectatic regions might explain why $R_n$ was increased after VILI during $D_{\text{rec}}^{\text{PEEP}}$. However, these changes in $R_n$ were relatively minor, and were even non-existent during $D_{\text{rec}}^{\text{PEEP}}$, indicating that central airway structure was not substantially affected by injurious ventilation. Indeed, stereological analysis revealed only occasional edema in the airways, and we did not find any airways that were either partially buckled or fully collapsed.

The structural and functional changes detailed above are associated with changes to the alveolar fluid-mechanical microenvironment that oppose alveolar inflation and promote collapse. The increased volume of fluid (Figure 5F) in partially flooded alveoli decreases the interfacial radius of curvature, which increases the pressure differential across the air–liquid interface and causes the lining fluid to exert a greater inward force that facilitates collapse. Furthermore, the functionality of the surfactant system is compromised (Figure 8C) from an apparent increase in surfactant degradation (increased SA fraction and decreased LA/SA ratio) and inactivation of PLs in the LA aggregates. The degradation of surface-active LA (Yamada et al., 1990) into surface-inactive SA (Figure 8A) may be attributed to

### Table 1: Phospholipid Content

| Condition   | Amount (nmol) | Total Phospholipid | LA | SA |
|-------------|---------------|--------------------|----|----|
| Control, PEEP=0 | 300           | 700                | 100| 200|
| Control, PEEP=3 | 400           | 900                | 150| 250|
| Control, PEEP=6 | 500           | 1100               | 200| 300|
| VILI, PEEP=0     | 600           | 1200               | 250| 350|
| VILI, PEEP=3     | 700           | 1400               | 300| 400|
| VILI, PEEP=6     | 800           | 1600               | 350| 450|

### Figure 7: Pulmonary System Elastance $H$ at the End of $D_{\text{rec}}^{\text{PEEP}}$

![Pulmonary system elastance graph](image)

### Figure 8: Total Phospholipid Content

![Phospholipid content graph](image)

**A** Total phospholipid content (white bar) is increased following ventilation and was accompanied by an increase in small aggregate (SA) and slight decrease in large aggregate (LA) causing a decrease in the LA/SA ratio (B). The total protein content in the BAL (C) is significantly increased in the VILI animals and the minimum surface tension measured using the pulsating bubble surfactometer (PBS) was significantly higher post-injury. Significant differences are indicated with asterisks and individual datapoints representing the pooled analysis of three animals are indicated with circles. BALF from three mice was pooled for each of the three datapoints. Error bars show standard error.
large cyclic strains imposed by high-volume, PEEP = 0 cmH\textsubscript{2}O ventilation (Greenfield et al., 1964) that severely compresses the interfacial film so that it ruptures on re-expansion (Wyszogrodski et al., 1975). These mechanical effects can occur in the first 5 min of high tidal-volume ventilation (McClenahan and Urtnowski, 1967; Forrest, 1972). Therefore, in the present study, ventilation-induced surfactant degradation may have been the first step in the vicious cycle of altered alveolar dynamics, collapse, atelectrauma, and volutrauma that lead to eventually fatal VILI.

The mechanism for inactivation of the LA fraction is less clear and may very well be multifactorial. Airspace protein content is elevated (Figure 8C) due to alveolar surface damage (Dreyfuss et al., 1988; Hamlington et al., 2018b) and the resulting alveolo-capillary leak (Hamlington et al., 2018a). Alveolar surface damage increases the BALF concentration of epithelial proteins such as E-cadherin while alveolo-capillary leak accounts for the presence of serum proteins including albumin and immunoglobulin (IgG) in the BALF of mice ventilated under similar conditions (Smith et al., 2017). Our surface tension analysis includes the effects of BAL proteins that migrate with the PLs of the LA fraction during ultracentrifugation and inhibit surface lowering activity through disruption of phospholipid packing in the molecular films at the air–liquid interface. Furthermore, changes to the LA composition including phospholipids, surfactant-associate proteins or both may be present, and were not explored in these studies. Preliminary studies examining the presence of secretory phospholipases capable of targeting surfactant PLs revealed no significant changes in the BAL during this VILI model (data not shown) (Hite et al., 1998).

Our study has a number of limitations. First, the lungs were prepared for morphometric analysis with the chest wall retracted, which means that at high inflation pressures the lung volumes would have been greater than in closed-chest animals due to absence of constraint by the thorax. Conversely, at low inflation pressures the absence of the chest wall would have resulted in lower lung volumes due to the absence of the negative pleural pressure that opposes collapse. Based on lung volumes, airway pressures, and esophageal pressures measured in mice of equivalent age and weight to those used in the current study (Lai and Chou, 2000) we estimate that our inflation pressures the absence of the chest wall would have been greater than in closed-chest animals and inhibit surface lowering activity through disruption of phospholipid packing in the molecular films at the air–liquid interface. Furthermore, changes to the LA composition including phospholipids, surfactant-associate proteins or both may be present, and were not explored in these studies. Preliminary studies examining the presence of secretory phospholipases capable of targeting surfactant PLs revealed no significant changes in the BAL during this VILI model (data not shown) (Hite et al., 1998).

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**CONCLUSION**

Three alveolar phenotypes typically develop with VILI: (1) flooded alveoli that cannot be recruited at any pressure, (2) unstable alveoli that are open at high pressures but readily collapse as pressure is reduced, and (3) relatively normal alveoli that remain open at low pressures. The relative proportions of each phenotype can be estimated using derecruitability tests that quantify time-dependent alveolar derecruitment as a function of PEEP. These correlations between lung structure and function could provide insights into optimally lung-protective ventilation strategies.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study are available on request to the corresponding author.

**ETHICS STATEMENT**

The animal study was reviewed and approved by University of Vermont, Institutional Animal Care and Use Committee and the University of Colorado Denver Anschutz Medical Campus, Institutional Animal Care and Use Committee.

**AUTHOR CONTRIBUTIONS**

BS, GR, KH, LK, MO, RH, and JB contributed to the conception and design of the study. BS, GR, AC, CM, KO, CC, and MN acquired the experimental data. BS, KH, RH, and JB analyzed and interpreted the data. BS drafted the manuscript. CM, KO, RH, and JB wrote sections of the manuscript. All authors contributed to the article and approved the submitted version.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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