Real Time Lung Imaging for the Detection of Lung Injury and Alveolar Fluid Movement During Mechanical Ventilation

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Abstract: Experimental ventilator-induced lung injury (VILI) is characterized by alterations in alveolar epithelial and microvascular permeability that favors the systemic dissemination of lung borne cytokines or bacteria. Animal models of VILI have been shown relevant to patient care and outcome and help explaining why most patients with the acute respiratory distress syndrome do not die from respiratory failure but from multiple organ dysfunction. Recent experimental studies also showed that adverse ventilator patterns may propel airway secretions and bacteria to previously healthy lung regions. Noninvasive imaging techniques were used for years to study the net rate of protein flow across the pulmonary microvascular endothelium and the alveolar epithelium

INTRODUCTION

Gas is separated in the alveolar spaces from blood by a thin membrane composed of epithelial (type 1 alveolar cells) and endothelial cell layers that are either closely juxtaposed or separated by a small interstitial space. The alveolocapillary barrier is thin enough to allow efficient gas exchange and strong enough to prevent the burst of plasma proteins and water into alveoli. This barrier may however experience physical, chemical or biological insults that lead to an increased permeability to proteins [1-5]. Because the permeability of the alveolo-capillary membrane to water is much greater than that to proteins, the measurement of extravascular lung water is rather insensitive for detecting pulmonary edema of the permeability type. Imaging techniques providing a gross estimation of the epithelial and the endothelial layer permeability to proteins have been developed for more than 40 years and have had, physiological, pharmacological and clinical applications [6]. Depending of its route of administration, via the airways [7-9] or intravenously [10-12], a radiolabeled protein can be used for estimating epithelial or microvascular permeability, respectively.

Keywords: Radionuclide imaging, respiration, artificial, blood-air barrier, acute lung injury.

VENTILATOR-INDUCED LUNG INJURY

The deleterious effects of mechanical ventilation on lungs have been for long referred to as barotrauma. For many years, clinicians defined barotrauma as the occurrence of air leaks resulting in the accumulation of extra-alveolar air responsible for a number of manifestations, of which the most threatening is tension pneumothorax [15]. In addition to these "macroscopic" events, whose adverse consequences are usually immediately obvious, mechanical ventilation may produce more subtle physiological and morphological alterations, especially when high airway pressures, and thus high tidal volumes (VT), are used. Unlike the classic forms of barotrauma, our knowledge of these alterations has come only from experimental studies. Alterations in lung fluid balance, increases in endothelial and epithelial permeability,
and severe tissue damage have been seen following mechanical ventilation in animals [16]. The clinical relevance of VILI was highlighted by the Acute Respiratory Distress Syndrome (ARDS) Network trial [17] that showed 22% reduction of mortality in patients with ARDS when the mechanical stress applied to the lungs was lessened by a reduction in V_T. On a mechanical point of view, four specific VILI mechanisms have been identified: a) regional overdistension caused by the application of a local stress or pressure that forces cells and tissues to assume shapes and dimensions that they do not assume during unassisted breathing [13, 14, 18]; b) so-called “low volume injury” associated with the repeated recruitment and derecruitment (airway collapse and reopening, liquid movement in distal airways) of lung units, which causes the abrasion of the epithelial airspace lining by interfacial forces [19, 20]; c) the inactivation of surfactant triggered by large alveolar surface area oscillations [21, 22]; d) interdependence mechanisms that raise cell and tissue stress between neighboring structures with differing mechanical properties [23].

Lung inflation resulting in too high tissue stretch also promotes the systemic dissemination of lung borne cytokines [24, 25] or bacteria [26-28]. Although these observations may offer an explanation for the observation that most ARDS patients do not die from respiratory failure but from multiple organ dysfunction syndrome [29], similar experiments yielded conflicting findings [30, 31], thus questioning the exact relationship between VILI and P. This finding led to the suggestion that “translocating” bacteria came from the alveolar spaces. In addition, animals ventilated with a large V_T and a low positive end-expiratory pressure (airway collapse and reopening, liquid movement in distal airways) of lung units, which causes the abrasion of the epithelial airspace lining by interfacial forces [19, 20]; c) the inactivation of surfactant triggered by large alveolar surface area oscillations [21, 22]; d) interdependence mechanisms that raise cell and tissue stress between neighboring structures with differing mechanical properties [23].

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Imaging of Alveolar Epithelial Permeability

Biophysical Basis

This technique involves the administration, by aerosolization or instillation, of a radioactively labeled tracer in the airways. The clearance of tracer activity from the lung is then measured, usually with some external radiation detection system such as small radiation probes or larger gamma cameras. The decrease of thoracic activity over time is a function of the following variables: i) lipid solubility will determine the ability of the tracer to dissolve in and then diffuse across lipid cellular membranes. This property is shared by gases, alcohol and other lipophilic solutes which penetrate these barriers so quickly that equilibration between fluid injected into the alveoli and the blood is nearly complete during the capillary transit time (0.75 s) [35]. In contrast, lipophobic solutes such as the electrolytes, sugars and proteins are largely confined to extracellular pathways and require no less time than several minutes before concentrations in the air spaces start to decline. ii) The molecular weight of the tracer is inversely correlated with its rate of diffusion throughout the alveolo-capillary barrier [36]. iii) The permeability surface area product of the alveolar epithelium. As previously published [37], the relationship between the alveolar concentration of the tracer (C/C_0) and the permeability (P) surface (S) area product over time (t) is as follows (see appendix):

\[ C/C_0 = e^{(PS/V)t} \]

where \( C_0 \) is the initial indicator concentration, C the indicator concentration, and V the volume of fluid instilled in the airways [37]. The slope of C/C_0 on a logarithmic coordinate against linear time would then be the constant – PS/V. Distention of the lung if alveoli open by unfolding will tend to increase V more than S [37]. One might therefore expect S/V and the value of PS/V to decrease with lung inflation. In fact, the decline in indicator concentration is accelerated by distension in instillation experiments, suggesting an increase of P. This finding led to the equivalent pore approach (i.e., the stretching of interendothelial pores) used by Egan [38, 39].

Practical Aspects

Beyond the above-mentioned variables, other factors can impact on the thoracic clearance of a tracer administered in the airways: i) heterogeneous spread of the tracer in the lungs due to regional changes in ventilation and, sometimes, inability to ensure adequate delivery of the tracer to the most injured regions (and thus, often the most poorly ventilated); ii) uncertainty about the site of clearance (e.g., airway vs alveolus); iii) variability of the radiochemical purity of the tracer (i.e., the proportion of the total activity of the isotope bound to its carrying protein or molecule). An ideal tracer would be tightly bound to the isotope, it would be relatively simple to obtain and prepare, and its clearance rate would be fast enough to allow accurate measurements in a reasonable length of time [8, 40]. In addition, an increase in its clearance rate would be uniquely associated with lung injury and not lung inflation.
Peterson et al. considered these issues and compared the clearance rates of aerosolized 99mTc-labeled diethylenetriaminepentaacetate (99mTc-DTPA, mol wt 492 Da), 99mTc-labeled albumin (mol wt 69 000 Da) and 99mTc-aggregated albumin (mol wt 383 000 Da) from the lungs of anesthetized sheep subjected to lung inflation or lung injury induced by intravenous oleic acid [41]. Spontaneous clearance of 99mTc-DTPA was high (0.42±0.15%/min) and significantly and equivalently increased after oleic acid injection and application of a PEEP of 10 cmH2O. 99mTc-albumin exhibited a much lower spontaneous clearance (0.06±0.02%/min) that was not modified by PEEP application and increased after oleic acid injection. The 99mTc-aggregated albumin clearance rate was unchanged by lung inflation and increased slightly with lung injury. The authors concluded that 99mTc-albumin was the best indicator for studying the effects of lung epithelial injury on protein transport out of the air spaces.

**Imaging of Pulmonary Microvascular Permeability**

Measuring the time-dependent accumulation of an intravenously administered radioactively labeled protein tracer into lung tissue (i.e., from plasma to interstitium) has been used for years to determine whether or not the integrity of the capillary endothelial barrier was compromised [6, 42]. However, since the tracer moves to other tissues, its plasma concentration decreases over time. Therefore, the count rate decreases at the detector, which might overwhelm the slight increase due to lung influx of the tracer. Normalization of the activity of the tracer is thus required. In 1957 Aviado and Schmidt attempted to follow the development of alloxan-induced pulmonary edema using diffusible (131I-albumin) and nondiffusible (32P-labeled red blood cells) tracers [43]. Due to technological limitations, they could only use one tracer in each animal. They however observed and increase in 131I counts in lungs occurring at a time when 32P counts were known to decrease. They suggested that this reflected increased extravasation of plasma albumin in lung parenchyma. In 1968, Potchen and Welch successfully distinguished gamma emissions from simultaneously injected diffusible (131I-albumin) and nondiffusible (32P-labeled red blood cells) tracers using a pulse height analyzer [44] thus allowing larger applications of this technique. Gorin et al. further validated the physiological relevance of this method by showing a correlation between the pulmonary 111In-transferrin transvascular flux and the tracer protein accumulation in lung lymph in sheep submitted to various types of lung injury [45]. The same authors showed the feasibility of this technique in humans, although the need to label red blood cells in vitro before restituting them to the patient remained a major constraint [46]. Rosell et al. finally simplified and homogenized this technique by validating the normalized slope index, that is, “the time rate of change of radioactivity originating from protein in lung interstitium divided by this ratio at time 0, and corrected for blood volume changes”. Only one tracer could be used providing the decrease of its plasma activity over time would be accounted for by measuring the activity of a fixed plasma volume at different time points and then normalizing the detector count rate by the plasma count rate [47, 48] or by selecting a vascular region with a gamma camera in order to quantify plasma protein clearance without the need for a blood sample [49-51].

Indium (113mIn or 111In) has been widely used as it strongly binds to transferrin in vivo and forms a good tracer of passive protein movement because it does not bind to transferrin receptors [52]. Basran et al. noninvasively measured its pulmonary accumulation in ARDS patients at bedside using a portable probe radiation detector. They showed clearly separated values of accumulation indexes between ARDS patients and healthy subjects [10]. They further used this method to evaluate the effect of a pharmacological intervention on pulmonary microvascular permeability. They observed that patients whose pulmonary protein accumulation indexes decreased after terbutaline challenge (i.e., responders) were more likely to survive than those who did not respond [53]. The growing enthusiasm for this technique was however reduced when Schuster et al. failed to discriminate patients with noncardiogenic and cardiogenic pulmonary edema using the normalized slope index after intravenous injection of 99mTc-albumin [54]. The authors hypothesized that the overlap observed in the permeability indexes of the two groups of patients was due to the occurrence of capillary stress failure associated with high intravascular pressures which may have allowed rapid egress of radiolabeled proteins from the vascular space [55].

**Simultaneous Imaging of Alveolar Epithelial and Pulmonary Microvascular Permeability to Proteins**

**Physiological Relevance**

Permeability to proteins of the epithelial and endothelial layers widely differs, alveolar epithelium being much less permeable than the endothelium [11]. Moreover, the response of each layer of the alveolo-capillary membrane to physical, chemical or biological insults may differ, depending on insult type [2, 4, 5]. For instance, *Pseudomonas aeruginosa* elastase on the alveolar side produces alveolar epithelium leakiness that manifests as an increase in the clearance of aerosolized albumin, without evidence of edema [1], whereas chronic heart failure increases protein fluxes in both directions [56]. During lung inflation, alveolar permeability to proteins has been shown to increase in sheep [41, 57] and in humans [58]. On the other hand, short periods of overinflation resulted in a reversible increase in pulmonary microvascular permeability to albumin and perhaps some increase in alveolar epithelial permeability as radiolabeled albumin injected in the systemic circulation was recovered in bronchoalveolar lavage fluid [59]. More sustained high tidal volume ventilation produced endothelial and epithelial cell alterations and a pulmonary edema of the permeability type [16]. Noninvasively monitoring the simultaneous changes of lung microvascular and alveolar permeability to proteins in vivo with a model combining an alveolar and a systemic tracer allowed us to independently examine the response of each layer of the alveolo-capillary barrier during lung inflation [60].

**Description of the Double-Isotope Model**

Briefly (see reference [60] for a more detailed description), male Wistar rats were anesthetized, tracheostomized and mechanically ventilated. About 400 μCi of 111In chloride were injected intravenously. A solution...
containing $^{99m}$Tc-albumin (~600 μCi) was selectively instilled in one lung. Pulmonary microvascular permeability was estimated using the normalized lung-to-heart activity ratio of $^{111}$In-transferrin which is related to the normalized slope index [61]. This index (lung-to-heart ratio divided by its initial value) accurately reflects $^{111}$In-transferrin plasma to lung flux assuming that lung and heart blood volumes do not vary during the observation period. Alveolar epithelial permeability was estimated by measuring the decrease of $^{99m}$Tc-albumin in a region of interest (ROI) drawn over the thorax and fitted with a two-exponential decay equation. Scintigraphic imaging was performed as follows: continuous planar thoracic acquisitions were performed using a small animal gamma camera. Acquisition windows were 140 KeV±15% for $^{99m}$Tc and 245 KeV±20% for $^{111}$In ($^{99m}$Tc does not emit in this energy window). Spillover of $^{111}$In in the $^{99m}$Tc energy window (~60% $^{111}$In counts in the 245-KeV window) was subtracted, and $^{99m}$Tc activity decay was taken into account. $^{99m}$Tc/$^{111}$In lung count ratio in the $^{99m}$Tc window was always ~10 so that the correction always remained below 10% of $^{99m}$Tc counts. Acquisition lasted 120 min without interruption. A 30 min long quality control acquisition was performed at the beginning of each experiment while ventilating the animal with protective settings in order to make sure that the baseline diffusion of each tracer through the alveolo-capillary barrier was negligible. An intervention could be performed once this acquisition was completed.

**Applications of the Model**

We assessed the effect of lung inflation on alveolar epithelial and endothelial permeability to proteins by ventilating 4 groups of rats with different end-inspiratory plateau pressures (Pplat). Pplat tested were 15, 20, 25 and 30 cmH$_2$O, corresponding to V$_T$ of 7.8±0.31, 13.7±4.69, 22.2±2.12 and 25.9±1.76 mL/kg. The main finding of this study was that the same end-inspiratory pressure threshold (between 20 and 25 cmH$_2$O) was observed for epithelial and endothelial permeability changes (Fig. 1). Interestingly, this threshold pressure approximately corresponds to the pressure at which a decrease in the respiratory system pressure-volume curve slope (the so-called “upper inflection point”) is observed in rats [62], which may reflect the occurrence of lung tissue overstretch. Whereas protein flux from plasma to alveolar space was constant over time, the rate at which $^{99m}$Tc-albumin left air spaces decreased with time, as previously described [41, 63]. Application of a three-compartment model including an intermediate interstitial space whose volume could change over time explained this pattern. This simple double-isotope imaging technique allowed to noninvasively explore the simultaneous changes of lung microvascular and alveolar permeability to proteins in vivo.

This model was further used to assess the effect of pharmacological interventions in the setting of VILI. Changes in epithelial and endothelial permeability induced by injurious mechanical ventilation with or without PEEP were challenged by a β$_2$-agonist (terbutaline) administered either intratracheally or intraperitoneally [64]. High volume ventilation resulted in immediate leakage of $^{99m}$Tc-albumin from alveolar spaces and increased pulmonary uptake of systemic $^{111}$In-transferrin. Terbutaline in the instilled solution and PEEP lessened alveolar $^{99m}$Tc-albumin leakage and $^{111}$In-transferrin uptake due to high volume ventilation, whereas terbutaline given intra-peritoneally only lessened $^{111}$In-transferrin uptake. Terbutaline in the instilled solution also lessened the increase in lung wet-to-dry weight ratio due to high volume ventilation. Our simple double-isotope imaging model allowed us to first describe the effect of a β$_2$-agonist on alveolar and endothelial permeability changes during high volume ventilation. The effect of PEEP was likely due to a decrease in tidal volume excursions, because it has been shown that beyond absolute stretch level, the magnitude of a cyclic stretch also affected cell response to mechanical challenge [65].

**NONINVASIVE ASSESSMENT OF INTRAPULMONARY FLUID DISPERSION DURING MECHANICAL VENTILATION**

As seen above, mechanical ventilation may favor systemic dissemination of sepsis and inflammation during...
Imaging VILI

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Fig. (2). Examples of scintigraphy images integrating the 15 min following instillation (t0-t15; left panels) and the last 15 min of the experiment (t195-t210; right panels). Regions of interest (ROIs) were drawn around initial focus of edema (ROI_E), the apex of the same lung (ROI_A), the contralateral lung (ROI_CL) and over the thorax (ROI_T). At baseline (left panels), all animals were ventilated with a tidal volume of 8 mL/kg and a PEEP of 2 cmH_2O and exhibited focalized localization of the tracer in the left lung. When the same ventilator settings were kept during the experiment (a), the tracer remained remarkably confined in the initial zone; there was no contra-lateral and slight homolateral dissemination. High volume ventilation (Pplat=30 cmH_2O) with no PEEP (b) induced strong homo and contralateral dispersion of the tracer and systemic leakage as attested by the evident decrease in overall activity. High volume ventilation with 6 cmH2O PEEP (c) induced systemic, but not contralateral, dissemination of the tracer. Reproduced from [66] with authorization.

Fig. (3). Activity changes in the four ROIs (as defined in the legend of Fig. 2) expressed as percentage of initial total activity for 6 animals ventilated with a tidal volume of 8 mL/kg and a PEEP of 2 cmH_2O for the 30 first min. Notice that the activity is remarkably stable in all ROIs. When high volume ventilation with no PEEP was started (after 30 min), there was an almost immediate decrease of activity in ROI_E and ROI_T and a dramatic increase in ROI_CL. Changes in ROI_T activity displayed a two phase’s exponential decay: activity decreased fastly between t30 and t60 and more slowly between t60 and t210. Reproduced from [66] with authorization.

Bacterial pneumonia. There is also growing evidence that ventilator settings may localize or disperse proteinaceous lung edema or bacteria. In an unilateral model of Pseudomonas aeruginosa pneumonia in rats, Schortgen et al. showed that high volume ventilation with no PEEP (b) induced strong homo and contralateral dispersion of the tracer and systemic leakage as attested by the evident decrease in overall activity. High volume ventilation with 6 cmH2O PEEP (c) induced systemic, but not contralateral, dissemination of the tracer. Reproduced from [66] with authorization.

disperse localized radiolabeled alveolar edema to the opposite lung [66]. A 99mTc-labeled albumin solution was instilled in a distal airway. This protocol produced a zone of alveolar flooding that stayed localized during conventional ventilation or spontaneous breathing. High end-inspiratory pressure ventilation dispersed alveolar liquid in the lungs. This dispersion was prevented by PEEP even when VT was the same and thus end-inspiratory pressure even higher (Fig. 2). Interestingly, contralateral liquid dispersion began almost immediately after high volume ventilation was started, suggesting that this dispersion might be the consequence of a convective movement induced by the ventilation (Fig. 3). High-inspiratory and -expiratory flows favored fluid transport back and forth toward the alveoli and the airways. Moreover, in the heterogeneous lung, fluid transfer may be propelled toward regions of normal lung because they have a normal compliance and thus receive high inspiratory flows. PEEP may have prevented dispersion by avoiding lung collapse and stabilizing edema fluid in the distal airways. These findings are in keeping with and help understanding the intrapulmonary dispersion of P. aeruginosa during a similar ventilation modality [28]. The clinical relevance of the movement of such noxious biofluids in the airways has recently been addressed [67].

CONCLUSION

Simple noninvasive imaging techniques can be used to noninvasively study the changes in lung microvascular and alveolar permeability to proteins in vivo and the movement of fluids in the lungs during mechanical ventilation. Continuously and simultaneously monitoring such variables might be of great interest in experimental research on VILI as it allows the assessment of physiological or pharmacological interventions [68, 69].

APPENDIX

Relationship between changes in the alveolar concentration of the tracer (C/C_0) and the permeability surface area product of the alveolar epithelium: as previously
published [37], the rate of solute diffusion (dQ/dt) through membranes is directly proportional to membrane surface area (S) and the concentration gradient (dc/dx) in accordance with Fick’s First Law of Diffusion:
\[
dQ/dt = -DS \, dc/dx
\]  
(2)
where D represents the diffusion coefficient per unit of surface area. As the concentration difference across the membrane Δc can only be measured, this equation becomes:
\[
dQ/dt = -PS \Delta c
\]  
(3)
where PS is the whole membrane permeability (P) surface (S) area product. Assuming that much of the fluid instilled into the airways will reach alveoli, the indicator concentration (C) of the fluid will decline uniformly in the volume (V) instilled:
\[
dQ/dt = V \, dC/dt = -PS \Delta c
\]  
(4)
If plasma concentration remains low, this relationship can be simplified to:
\[
dC/dt = -PSC/V
\]  
(5)
Solution of equation (4) yields the following:
\[
C/C_0 = e^{-PS/V}t
\]  
(6)
where C0 is the initial indicator concentration [37].

REFERENCES
