Protecting the lungs but hurting the kidneys: causal inference study for the risk of ventilation-induced kidney injury in ARDS

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The acute respiratory distress syndrome (ARDS) is the most serious form of acute respiratory failure in the intensive care unit (ICU), defined by hypoxemia and bilateral opacities on chest radiographic imaging (1). ARDS can represent the end-result of many different insults—such as sepsis, pneumonia, aspiration or pancreatitis—leading to injury of the alveolar epithelial/endothelial interface with resultant permeability edema (2). Epidemiologically, ARDS accounts for up 10% of all ICU admissions and can be diagnosed in up to 23% of mechanically-ventilated patients (2). Severe ARDS carries 30–40% mortality risk, whereas ARDS survivors often suffer from chronic critical illness and long-term neurocognitive deficits (3). Despite advancements in understanding ARDS pathogenesis, no pharmacologic therapy for ARDS has shown efficacy in clinical trials. Thus, invasive mechanical ventilation (IMV) remains the mainstay of ARDS management, with the goals of supporting gas exchange, allowing the respiratory system to “rest”, and providing time to treat the underlying pathology.

Decades of research have shown that despite its beneficial effects in gas exchange, IMV can also precipitate and perpetuate severe lung injury. Ventilation-induced lung injury (VILI) encompasses a spectrum of mechanisms that relate to the delivery of positive-pressure ventilation in lungs with low compliance and heterogeneously distributed pathology (i.e., worse edema in dorsal lung units and small amounts of normally aerated lung ventrally). Over-distention of the normally aerated alveoli due to inappropriately high tidal volumes results in stretch injury (volutrauma), with further risks of alveolar rupture and air leak (e.g., pneumothorax) (4). Conversely, ventilation at inappropriately low volumes leads to cyclic alveolar opening and collapse, with ensuing repetitive epithelial injury from shear forces (atelectrauma). Such mechanisms converge into the release of inflammatory mediators from the injured lung (e.g., interleukin-6 and tumor necrosis factor-alpha), which can leak in the systemic circulation due to the increased alveolar/capillary permeability and cause distant end-organ damage (biotrauma) (5). To mitigate VILI, multiple lung protective ventilatory strategies have been examined, yet only two elements have consistently proven to be important: low tidal volume ventilation (typically prescribed at 6 mL/kg of ideal body weight) to prevent volutrauma (6) and the application of positive end-expiratory pressure (PEEP) to prevent atelectrauma (7). Nonetheless, delivering an optimal PEEP individualized to patient-specific respiratory physiology remains an issue of ongoing debate (7).

While lung protective ventilation remains the central tenet of ARDS management, the extrapulmonary consequences of IMV cannot be overlooked. Patients with ARDS often develop cardiovascular, renal and neurologic dysfunctions, and then succumb to consequences of
multiple organ failure syndrome rather than to refractory hypoxemia. Acute kidney injury (AKI) is the most frequent extrapulmonary organ damage in ARDS and affects up to 50% of patients, doubling their mortality risk compared to patients without AKI (8,9). Initiation of IMV is associated with a threefold increase in the odds of AKI, a finding consistent among populations of critically-ill patients, regardless of ARDS diagnosis (10). Such epidemiologic observations fueled interest in understanding the mechanisms involved in the cross-talk between the lungs and the kidneys that can lead to ventilator-induced kidney injury (VIKI) (11).

Data from clinical studies and physiologic experiments for plausible lung-kidney interactions during IMV can be synthesized into three main mechanisms (Figure 1). First, gas exchange disturbances from the primary lung pathology can exert direct effects to the kidneys. The hallmark hypoxemia of ARDS and permissive hypercapnia can alter renal vascular resistance and perfusion pressures, resulting in a drop of glomerular filtration rate (GFR), ischemic acute tubular necrosis and finally, clinical AKI (Figure 1A) (11,12). Second, the kidneys are directly exposed to the hemodynamic effects of IMV on the systemic circulation. Positive pressure ventilation increases the intrathoracic and right atrial pressures, impairing right ventricular venous return and leading to renal venous congestion. Apart from the negative effects on right ventricular preload, intrathoracic pressures also raise the right ventricular afterload, with a net reduction of cardiac output. As recipients of a diminished cardiac output, the kidneys are exposed to the double hit of lower renal blood flow coupled with venous congestion, ultimately lowering GFR (Figure 1B). Finally, renal glomeruli and tubules are also exposed to the inflammatory mediators released from the "leaky" injured lungs of ARDS (biotrauma), which can further disturb intra-renal hemodynamics, vascular tone and endothelial/epithelial integrity (Figure 1C) (8). The relative contributions of these purported mechanisms in generation of VIKI have not been clarified, yet these are important for the design and implementation of kidney protective ventilatory strategies.

In the last issue of the Annals of Translational Medicine, Leite et al. addressed the important question of which parameters of mechanical ventilation parameters directly influence the risk of VIKI, by doing causal inference analyses in an observational cohort study (13). Starting with a large, open-access database (Multiparameter Intelligent Monitoring
The mechanisms of lung-kidney cross-talk during IMV are intertwined and can translate into a dataset of multiple correlated and mathematically-derived variables. Therefore, the authors used a causal modeling approach with directed acyclic graphs (DAGs) to ascertain which of the statistical associations may represent true causal risk factors for VIKI. DAGs represent a method of increasing popularity in causal inference studies, due to their intuitive visual representation of data and interpretability of results. In contrast to studies examining the use of a set of variables to predict an outcome of interest (i.e., predictive modeling), causal inference studies aim to estimate the direct “causal” effect of an exposure on an outcome, so that causal hypotheses can be generated to guide the design of clinical trials. By definition, a DAG consists of a set of vertices (variables) and a set of (directed) edges, representing conditional dependencies between variables. An edge between A→B implies that a change in A will “cause” a change in B, independently of all other variables or subsets of in the dataset (15). Using a graph to depict a model enables easy and succinct display of substantive assumptions collectively, simplifies the calculation of the joint probabilistic density function and also provides an intuitive visualization of causal relationship network among variables of interest. Nevertheless, DAGs have certain limitations including the assumption of linearity of interactions and potentially overlooking unobserved confounder variables, although progress has been made in these fronts as well (16).

Graphical models can be built by incorporating prior knowledge; or can be done as probabilistic graphical model (PGM) learning directly from observational data, through searching the space of potential graphs to maximizing an objective function (Bayesian Information Criterion, graph stability, etc.); or a combination of two approaches (17). Well-encapsulated tools like R packages of毡learn and psalq are used extensively when datasets contain a single type of variables, either continuous or discrete. As for causal discovery and PGMs, the widely acknowledged causal discovery software TETRAD now can learn graphs over mixed variable types (continuous and discrete data in the same dataset). The tool of graphical modeling is useful in biomedical research when modeling multi-scale data such as identifying genetic variants related to disease onset or response to drugs (18), or building predictive models for disease progression (19,20). In this case of VIKI for example, we are not just interested in whether levels of PEEP would help evaluate risk of VIKI adjusted for all other variables, but also whether high PEEP levels can actually cause VIKI so that then we could titrate PEEP appropriately to reduce the risk of VIKI. For that purpose, building a graphical model representing presumptions collectively would be helpful for analysis of causality.

In the study by Leite et al., the authors constructed a DAG by linking several IMV parameters (such as time-weighted PEEP, driving pressure, plateau pressure, tidal volume), gas exchange metrics (PaO\(_2\), PaCO\(_2\), and PaO\(_2\)/FiO\(_2\) ratio), as well as an overarching average respiratory system compliance (C\(_{rs}\)) with VIKI, while controlling for other clinical variables considered as confounders (e.g., age, gender, fluid balance, drugs etc.). Following a series of main and sensitivity analyses, two variables were consistently shown to have causal effects with VIKI: C\(_{rs}\) and PEEP. Each 5-mL/cmH\(_2\)O increase in C\(_{rs}\) reduced the odds of severe AKI by 10%, whereas each 1-cmH\(_2\)O increase in PEEP increased the odds of severe AKI by 5%.

The direct causal relationship of the average compliance of the entire respiratory system (C\(_{rs}\)) with VIKI is physiologically intuitive. This summary compliance (defined as change of volume over change of pressure with each mechanical breath, and thus measured as delivered tidal volume divided by the difference between the measured plateau pressure and PEEP) is a crude, global reflection of combined lung and chest wall mechanics. Low C\(_{rs}\) can thus represent the “stiff” and “wet” lungs of ARDS, as well as a restricted thoracic cavity, from obesity, effusions or increased intra-abdominal pressures. The authors reasonably asserted that the effects of C\(_{rs}\) on VIKI could be, at least partially, mediated by some of the measured respiratory and gas exchange variables. However, in mediation analyses, only PEEP was found to have a statistically significant (but minimal in size) effect on the C\(_{rs}\)-VIKI association. Thus, the demonstration of a causal effect for diminished C\(_{rs}\) underlines the complex cross-talk between lungs and kidneys, in that a global metric of respiratory dysfunction is linked to kidney dysfunction. However, this finding is not clearly actionable. Most evidence-based interventions in ARDS effectively aim to improve some of the C\(_{rs}\) components [e.g., lung protective ventilation to avoid
VILI, conservative fluid management to reduce lung edema and prone positioning to recruit dorsal alveolar units (2), therefore no clear target for further mitigating VIKI risk emerged from the causal diagram analysis.

On the other hand, the important causal association between PEEP and VIKI in the analysis by Leite et al. may offer opportunities for kidney protective adjustments of ventilatory settings. The near-linear relationship between rising PEEP levels and severe AKI, independent of plateau pressures, suggests that perhaps it is the continuously distending and not the cyclically applied pressure that may hurt the kidneys (i.e., the constant PEEP instead of the intermittently applied pressures from tidal volume delivery, Figure 2). The most biologically plausible explanation for the injurious effects of PEEP is through altering kidney hemodynamics (Figure 1B). Nonetheless, the authors were not able to demonstrate any significant mediation of available hemodynamics parameters (central venous or mean pulmonary arterial pressure) between high PEEP and VIKI. Whether this mediation analysis in a subset of the cohort failed to detect the hemodynamic consequences of PEEP due to small sample size or whether PEEP exerts its adverse effects on unmeasured hemodynamic (e.g., intra-abdominal pressure and renal venous congestion) or other parameters remains unknown. Converging evidence suggests that overzealous PEEP application to accomplish maximal alveolar recruitment (“open lung approach”) is hemodynamically detrimental and results in worse clinical outcomes (21,22). Ongoing research is trying to identify which patients may have a beneficial response to PEEP up-titration, defined either by improvements in gas exchange, lung mechanics or imaging markers (23). Emerging data suggest that complex, biological endotypes of ARDS (24,25) may offer improved insights in understanding treatment-effect heterogeneity and targeting our ventilatory (and other) interventions. This framework of predictive enrichment by ARDS biological endotypes may help us better model and understand the effects of IMV on extrapulmonary organ damage.

The importance of delivering patient-centric—rather than lung-centric—care in ARDS is self-evident. The study by Leite et al. reminds us of the knowledge gaps in trying to accomplish this well-shared goal. Supporting the failing lungs with evidence-based, lung-protective IMV settings may still have deleterious, yet underappreciated effects on extra-pulmonary organ functions, which ultimately, are the key determinants of outcome in ARDS. The statistical hits for causal effects of Cn and PEEP on the risk of VIKI are biologically plausible, and should be considered as hypothesis-generating for further prospective study; a validation within a larger, multi-center cohort as well as considering a PGM for causal discovery would certainly make such assertions more robust. Assembling the causality chain of VIKI and other organ injury in ARDS is necessary for refining IMV approaches from their current lung-protective to a broader, patient-protective paradigm.

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**Footnote**

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