Does the use of high PEEP levels prevent ventilator-induced lung injury?

O uso de níveis altos de PEEP previne a lesão pulmonar induzida pelo ventilador?

ABSTRACT

Overdistention and intratidal alveolar recruitment have been advocated as the main physical mechanisms responsible for ventilator-induced lung injury. Limiting tidal volume has a demonstrated survival benefit in patients with acute respiratory distress syndrome and is recognized as the cornerstone of protective ventilation. In contrast, the use of high positive end-expiratory pressure levels in clinical trials has yielded conflicting results and remains controversial. In the present review, we will discuss the benefits and limitations of the open lung approach and will discuss some recent experimental and clinical trials on the use of high versus low/moderate positive end-expiratory pressure levels. We will also distinguish dynamic (tidal volume) from static strain (positive end-expiratory pressure and mean airway pressure) and will discuss their roles in inducing ventilator-induced lung injury. High positive end-expiratory pressure strategies clearly decrease refractory hypoxemia in patients with acute respiratory distress syndrome, but they also increase static strain, which in turn may harm patients, especially those with lower levels of lung recruitability. In patients with severe respiratory failure, titrating positive end-expiratory pressure against the severity of hypoxemia, or providing it in a decremental fashion after a recruitment maneuver, is recommended. If high plateau, driving or mean airway pressures are observed, prone positioning or ultraprotective ventilation may be indicated to improve oxygenation without additional stress and strain in the lung.

Keywords: Acute respiratory distress syndrome; Ventilator-induced lung injury; Respiration, artificial

INTRODUCTION

Over the last few decades, several experimental and clinical studies have noted the relevance of physical mechanisms in generating or perpetuating ventilator-induced lung injury (VILI). Overdistention due to a high tidal volume (VT) or end inspiratory pressures, and the repeated opening and closing of distal bronchi and unstable alveoli resulting in high stress and strain, have been proposed as the main physical mechanisms responsible for VILI. The use of a low tidal volume instead of a large one led to a marked effect on survival in a large prospective, randomized, multicenter trial of patients with acute respiratory distress syndrome (ARDS), initiating the era of low tidal volume ventilation or protective ventilation. However, the use of high positive
end-expiratory pressure (PEEP) strategies has yielded conflicting clinical outcome results.

Positive end-expiratory pressure was been used to improve hypoxemia in patients with ARDS shortly after the first description of the syndrome.(3) Later, higher levels of PEEP along with recruitment maneuvers were proposed to prevent intratidal alveolar recruitment and improve survival. However, despite several translational and clinical studies, the effectiveness of these maneuvers remains controversial.

In the present article, we will present a short historical review on the use of high PEEP levels in patients with ARDS and will discuss some recent experimental and clinical trials in different clinical settings. In our view, the benefit from protective ventilation is mainly due to a decrease in stress and strain secondary to the use of a low tidal volume, and hence cyclic strain, in a highly heterogeneous lung. In contrast, the protective effect of PEEP on VILI is more debatable, as although it is highly effective at improving oxygenation, it may also increase strain and stress on the lung.

Lung injury at low lung volumes and the open lung approach

Ventilation that occurs at low lung volumes can cause injury through multiple mechanisms, including the repetitive opening and closing of airways and lung units, effects on surfactant function, and regional hypoxia.(1) Different experimental models have shown that the repetitive tidal recruitment and derecruitment (R/D) of small airways does occur at low or absent PEEP levels, promoting or increasing markers of VILI, while recruitment maneuvers and high PEEP levels result in improved oxygenation and less histological damage.

These observations are supported by two clinical trials using an open lung approach with high PEEP levels and low tidal volumes. These studies found positive results for this method when compared against a “conventional” strategy consisting of low to moderate PEEP and large tidal volumes.(4,5) The effect of PEEP in these studies should be assessed carefully, as tidal volume limitation in the open lung strategy could be responsible for the observed benefit.

The concept of “baby lung” and a pioneering study by Hickling on permissive hypercapnia(6) led several groups to conduct prospective studies that compared a tidal volume and/or pressure limitation strategy against a more conventional approach (Table 1).(2,4,5,7-9) The largest and most important of these studies showed that the use of a tidal volume of 6mL/kg IBW reduced mortality by approximately 25% compared with ventilation with 12mL/kg IBW in over 800 patients with ARDS.(2)

High PEEP strategies after the ARDSnet low tidal volume trial

After the ARDSnet low Vt study, three large randomized trials compared high and moderate PEEP strategies using low tidal volumes in both groups (Table 1).(10-12) None of these studies showed differences in mortality. However, a meta-analysis of these three studies suggested a small survival benefit from the high PEEP strategy in the subgroup of patients with a ratio arterial oxygen partial pressure to fractional inspired oxygen (PaO$_2$/FiO$_2$) < 200.(13) Considering only the studies from Meade et al.(8) and Mercat et al.,(9) which defined refractory hypoxemia $a$ priori, high PEEP strategies led to significantly fewer episodes of refractory hypoxemia and required fewer rescue therapies.(14)

A recent trial comparing an open lung approach (OLA study) with the ARDSnet study involved 200 patients with a PaO$_2$/FiO$_2$ ratio < 200 after a period of stabilization of at least 12 hours of protective ventilation, thus selecting a group with higher disease severity (Table 1).(15) This study had a low power to detect any relevant effect on mortality, but showed improved oxygenation and, more importantly, lower driving pressures, which may translate into lower dynamic strain (vide infra).(16,17)

A large randomized trial (ART) led by Brazilian investigators is assessing the effects of alveolar recruitment followed by decremental PEEP titration to optimize static compliance. This trial involving 1,100 patients is expected to be completed in 2017 and will provide important information on the effect of the open lung approach for patients with ARDS.(18)

Why were all these studies negative?

The use of PEEP makes sense for two reasons: first, by recruiting unstable alveoli, PEEP improves gas exchange and tissue oxygenation; second, PEEP reduces and redistributes the heterogeneous mechanical stresses of tidal ventilation.(19) Only the first assumption has proven to be true in patients, as the mechanical response to PEEP is highly variable in patients with ARDS.(20)
Animal experiments showing the benefits of high PEEP strategies usually use a highly recruitable model of lung damage, which does not necessarily translate to human ARDS. In contrast, most clinical trials in patients with ARDS have not assessed their recruitability (Table 1). Thus, the benefit of a high PEEP strategy in patients with severe ARDS and refractory hypoxemia may be obscured by the induction of overdistention and further lung injury in patients with less severe forms of respiratory failure, and thus less recruitable lungs.

An example of this lower level of recruitability occurs in the perioperative setting. A large clinical trial using a high level of PEEP (12cmH₂O) and recruitment maneuvers during open abdominal surgery showed no protection against postoperative pulmonary complications. In contrast, in 400 patients undergoing major abdominal surgery and at high risk of pulmonary complications, a strategy using a low tidal volume and moderate levels of PEEP decreased major pulmonary and extrapulmonary complications within the first 7 days, compared to a conventional strategy (Vt 10 - 12mL/kg IBW and no PEEP).

Global strain and cyclic strain

In a recent experimental model, Protti et al. demonstrated that a lung strain (the ratio between tidal volume and functional residual capacity) greater than 1.5 - 2 was necessary to induce lung damage in pigs without previous lung injury. In a second experiment, Protti et al. used several combinations of tidal volume (dynamic strain) and PEEP (static strain) to induce a similar level of global strain (the sum of static and dynamic strain) large enough to induce lung injury. Dynamic strain, also called cyclic strain, is mainly determined by tidal volume, while static strain represents the volume of gas caused by PEEP and may be well represented by mean airway pressure. A ventilatory strategy consisting of small dynamic (lower Vt) and large static (higher PEEP) strains decreased several markers of lung injury and mortality, suggesting that static strain is less harmful than dynamic strain.

In humans with ARDS, Caironi et al. showed that high PEEP levels decreased R/D only in patients with highly recruitable lungs, whereas no differences were observed in patients with lower levels of recruitability. However, strain increased with higher PEEP levels independent of lung recruitability. In a small set of patients with ARDS, we showed that global strain increased along PEEP levels and plateaued at airway pressure. More recently, increasing PEEP from 9 to 15cmH₂O along with low Vt ventilation did not decrease tidal R/D but consistently increased tidal recruitment and hyperinflation.

Lessons from high-frequency oscillatory ventilation clinical trials

High-frequency oscillatory ventilation (HFOV), by allowing greater end-expiratory lung volume while minimizing cyclic strain, resembles a high PEEP low Vt strategy, which seems ideal for lung protection in patients with ARDS. However, two recent multicenter, randomized trials did not show a survival benefit to this strategy, and in one study HFOV led to more deaths than a conventional approach. Mean airway pressure (Paw) in both HFOV arms was higher (above 25cmH₂O, Figure 1) than that of controls, which could reflect a higher global strain. As cyclic strain is minimized by HFOV (due to a much lower tidal volume), the higher global strain may only be a result of the higher static strain. The greater levels of vasopressor and intravenous fluid administration in the Oscillate trial, induced by a higher Paw, may help support this hypothesis.

In summary, in patients with moderate to severe ARDS, the higher global strain observed with HFOV may explain its lack of benefit –or even its harm– as found in recent trials, and may suggest a limit for PEEP titration. As high PEEP levels increase mean airway pressure, and hence static and global strain, Paw values above 25cmH₂O may suggest a limit when a more conservative prone or ultraprotective approach should be used.

Moving to ultraprotective ventilation

In contrast to the controversial data on PEEP, limiting tidal volume has been shown to be beneficial, leading to fewer complications and/or less mortality in different groups of patients with mechanical ventilation and becoming the standard for ventilation in critically ill patients. The negative results in recent trials of high versus low/moderate PEEP have been ascribed to the use of low Vt in both arms (along with moderate PEEP in controls), precluding the trigger for injurious ventilation.
Table 1 - Ventilatory parameters at 24 hours and mortality in clinical studies comparing a protective strategy, tidal volume (Vt) limitation, versus a control group (top panel); a strategy of high positive end-expiratory pressure versus low positive end-expiratory pressure or minimal distension (middle panel); and a conventional protective strategy versus high frequency oscillatory ventilation (HFOV) (lower panel) in patients with acute respiratory distress syndrome. The driving pressure of the respiratory system (ΔP) is calculated as the difference between the plateau pressure and positive end-expiratory pressure. Note that a larger difference of driving pressure between groups (Dif ΔP) is associated with differences in mortality.

| Author            | Year | N  | Vt   | P<sub>a</sub> | PEEP | ΔP | Mortality | Vt | P<sub>a</sub> | PEEP | ΔP | Mortality | Dif ΔP | p value<sup>†</sup> |
|-------------------|------|----|------|-------------|------|----|-----------|----|-------------|------|----|-----------|--------|---------------------|
| Brochard et al.  | 1998 | 108| 7.1  | 25.7        | 10.7 | 15 | 46.6      | 10.3| 31.7        | 10.7 | 21 | 37.9      | 6       | ns                  |
| Stewart et al.    | 1998 | 120| 7.2  | 22.3        | 8.6  | 13.7| 48.0      | 10.8| 26.8        | 7.2  | 19.6 | 46.0      | 5.9     | ns                  |
| Ranieri et al.<sup>‡</sup> | 1999 | 44 | 7.6  | 24.6        | 14.8 | * 9.8| 38.0      | 11.1| 31          | 6.5  | 24.5 | 58.0      | 14.7    | 0.19                |
| Brower et al.     | 1999 | 52 | 7.3  | 27          | 9.3  | 17.7| 50.0      | 10.2| 30          | 8.2  | 21.8 | 46.0      | 4.1     | ns                  |
| Amato et al.<sup>‡</sup> | 1998 | 53 | 6    | 31.8        | 16.3 | * 15.5| 38.0      | 12  | 34.4        | 6.9  | 27.5 | 71.0      | 12      | <0.001              |
| ARDSnet<sup>(33)</sup> | 2000 | 861| 6.1  | 25          | 9.4  | 15.6| 31.0      | 11.9| 33          | 8.6  | 24.4 | 39.8      | 8.8     | 0.007               |

**Note:**
- PEEP - positive end-expiratory pressure; Vt - tidal volume; ΔP - driving pressure; Dif ΔP - difference of driving pressure; HFOV - high frequency oscillatory ventilation; ns - not significant. * Ranieri and Amato studies also use high PEEP in the protective strategy. The p value refers to the differences in mortality between groups.

Recent data suggest that inhomoogeneity in human ARDS acts to increase stress and is associated with disease severity and mortality.<sup>(21,34)</sup> In an experimental model in pigs, applying very high stress and strain to the lung parenchyma leads to abnormal lung densities that are detected within 8 hours of ventilation at inhomogeneous interfaces and increase exponentially until lung edema develops after 20 hours.<sup>(35)</sup>

Independent of lung inhomogeneity and recruitable, tidal volume limitation will always suppress the main physical mechanisms involved in VILI. Using dynamic CT in nine patients with ARDS, lowering Vt from 12 to 6mL/kg IBW was found to not only decrease transpulmonary pressure and hyperinflation but also diminish the cyclic R/D of unstable alveoli.<sup>(36)</sup>

In a clinical setting, a small study of 10 patients with ARDS and plateau pressures of 28 - 30cmH<sub>2</sub>O despite a Vt of 6mL/kg IBW, a further decrease in Vt to 4mL/kg IBW and partial extracorporeal carbon dioxide removal reduced pulmonary cytokine concentrations after 72 hours.<sup>(37)</sup> The use of a Vt of 3mL/kg IBW along with extracorporeal CO<sub>2</sub> removal may have benefited patients with PaO<sub>2</sub>:FiO<sub>2</sub> ratios < 150, when compared with a Vt 6mL/kg IBW protective strategy.<sup>(38)</sup> Using dynamic CT, we showed that the reduction in Vt from 6 to 4mL/kg IBW decreased R/D, while partial pressure of carbon dioxide (PaCO<sub>2</sub>) and pH could be maintained at clinical levels if instrumental dead space was minimized.<sup>(39)</sup>
New evidence on protective ventilation in ARDS patients suggests that paralysis and prone positioning also have a major role in improving clinical outcomes.\(^{(40,41)}\) The striking data from these studies contrast with those comparing higher and lower PEEP settings. In particular, prone positioning may enhance the effects of high PEEP by preventing the negative effects of PEEP on tidal hyperinflation.\(^{(42)}\)

Summarizing these data, we suggest that the mechanical benefit of PEEP is most often found in patients with acute respiratory failure from 5 to 12 or 15cmH\(_2\)O, as alveolar recruitment prevails and oxygenation improves (Figure 2). At these PEEP levels, recruiting collapsed alveoli may also reduce driving pressure (dynamic strain), which could translate into less VILI.\(^{(15)}\) However, although there is no clear limit, the use of high PEEP levels above 12 or 15cmH\(_2\)O should be carefully titrated, as higher static strain and overdistention may prevail over recruitment.\(^{(28-30)}\)

In contrast, a decrease in tidal volume below physiological levels of 3 to 4mL/kg IBW will always confer the benefit of lower transpulmonary pressure, which is the main determinant of cyclic strain. Theoretically, a Vt of 0 should eliminate the cyclic R/D of unstable alveoli, but is accompanied by the constraints of hypercapnia and respiratory acidosis (Figure 3). This is the principle behind ultraprotective ventilation and extracorporeal membrane oxygenation. However, the role of these methods in severe respiratory failure has yet to be demonstrated.

![Figure 2](image1.jpg)

**Figure 2** - Effect of increasing levels of positive end-expiratory pressure on alveolar recruitment, tidal recruitment and derecruitment and static strain. From zero end-expiratory pressure to a positive end-expiratory pressure of 5cmH\(_2\)O, there was marked recruitment and a decrease in recruitment and derecruitment, which provided a protective effect. Positive end-expiratory pressure levels above 15cmH\(_2\)O should be carefully titrated, as the impact on recruitment is less evident and strain may increase.

![Figure 3](image2.jpg)

**Figure 3** - Effect of different tidal volumes on tidal recruitment and derecruitment, partial pressure of carbon dioxide levels and transpulmonary pressure. A decrease in tidal volume will always induce a decrease in transpulmonary pressure, but a very low tidal volume may increase partial pressure of carbon dioxide and decrease pH. Vt - tidal volume; R/D - tidal recruitment and derecruitment; PaCO\(_2\) - partial pressure of carbon dioxide; P\(_{L}\) - transpulmonary pressure.

**FINAL COMMENTS**

We strongly support the use of an open lung approach in patients with severe acute respiratory distress syndrome, as it decreases refractory hypoxemia.\(^{(13-15)}\) However, whether high levels of positive end-expiratory pressure prevent ventilator induced lung injury is still controversial. The clinical evidence suggests that tidal volume limitation is the cornerstone of protective ventilation. Thus, the proven benefit of high positive end-expiratory pressure strategies in decreasing refractory hypoxemia should be carefully weighed against the induction of added strain and overdistention, as it may be harmful under certain clinical conditions, such as in perioperative patients, patients with mild respiratory failure or patients with interstitial diseases.
Limiting tidal volume (and thus cyclic strain) and applying moderate positive end-expiratory pressure levels (between 8 to 12 cm H₂O) to prevent excessive stress and strain on the lung may be sufficient for most ventilated patients. In patients with severe respiratory failure, titrating positive end-expiratory pressure against the severity of hypoxemia or in a decremental fashion to obtain better compliance or driving pressure is recommended.¹⁵,¹⁷ When plateau pressures are above 30 - 35 cm H₂O, driving pressures are above 15 - 20 cm H₂O or mean airway pressures are above 25 cm H₂O, the adoption of prone positioning or ultraprotective ventilation may be indicated to improve oxygenation without inducing added stress and strain on the lung.

**RESUMO**

A distensão excessiva e o recrutamento alveolar pelo volume corrente foram defendidos como os principais mecanismos físicos responsáveis pela lesão pulmonar induzida pelo ventilador. A limitação do volume corrente demonstrou benefícios quanto à sobrevida naqueles com síndrome da angústia respiratória aguda e é reconhecida como a pedra fundamental da ventilação protetora. Em contraste, o uso de elevados níveis de pressão positiva expiratória final em estudos clínicos gerou resultados conflitantes e ainda é um assunto controvertido. Nesta revisão, discutimos os benefícios e as limitações da abordagem de pulmão aberto, e debatemos alguns recentes estudos experimentais e clínicos, referentes ao uso de níveis baixos e moderados de pressão positiva expiratória final. Também distinguimos o estiramento dinâmico (volume corrente) do estático (pressão expiratória final positiva e pressão média nas vias aéreas) e discutimos seus papéis na indução da lesão pulmonar induzida pela ventilação. As estratégias com elevada pressão positiva expiratória final claramente diminuem a hipoxemia refratária em pacientes com síndrome da angústia respiratória aguda, porém também aumentam o estiramento estático, que, por sua vez, pode ser lesiva aos pacientes, especialmente para aqueles com nível mais baixo de recrutabilidade pulmonar. Em pacientes com insuficiência respiratória grave, recomenda-se a titulação da pressão positiva expiratória final contra a gravidade da hipoxemia, ou sua aplicação de uma forma decrescente após manobra de recrutamento. Caso sejam observadas pressões de platô, driving pressure ou pressão média nas vias aéreas, a posição prona ou ventilação ultraprotetora podem ser indicadas para melhora da oxigenação, sem estresse adicional e estiramento dos pulmões.

**Descritores:** Síndrome do desconforto respiratório do adulto; Lesão pulmonar induzida por ventilação mecânica; Respiração artificial

**REFERENCES**

1. Slutsky AS, Ranieri VM. Ventilator-induced lung injury. N Engl J Med. 2013;369(22):2126-36. Erratum in N Engl J Med. 2014;370(17):1668-9.
2. Acute Respiratory Distress Syndrome Network, Brower RG, Matthay MA, Morris A, Schoenfeld D, Thompson BT, Wheeler A. Ventilation with lower tidal volumes as compared with traditional tidal volumes for acute lung injury and the acute respiratory distress syndrome. N Engl J Med. 2000;342(18):1301-8.
3. Bein T, Grasso S, Moerer O, Quintel M, Guerin C, Deja M, et al. The standard of care of patients with ARDS: ventilatory settings and rescue therapies for refractory hypoxemia. Intensive Care Med. 2016;42(5):699-711.
4. Amato MB, Barbos CS, Meideiros DM, Magaldi RB, Schettino GP, Lorenzi-Filho G, et al. Effect of a protective-ventilation strategy on mortality in the acute respiratory distress syndrome. N Engl J Med. 1998;338(6):347-54.
5. Ranieri VM, Suter PM, Tortorella C, De Tullio R, Dayer JM, Brienza A, et al. Effect of mechanical ventilation on inflammatory mediators in patients with acute respiratory distress syndrome: a randomized controlled trial. JAMA: 1999;282(1):54-61.
6. Gattinoni L, Pesenti A. The concept of “baby lung”. Intensive Care Med. 2005;31(6):776-84.
7. Brochard L, Roudot-Thoraval F, Roupie E, Delclaux C, Chastre J, Fernandez-Mondejar E, et al. Tidal volume reduction for prevention of ventilator-induced lung injury in acute respiratory distress syndrome. The Multicenter Trail Group on Tidal Volume reduction in ARDS. Am J Respir Crit Care Med. 1998;158(6):1831-8.
8. Stewar TE, Meece MD, Cook DJ, Granton JT, Hoddle RV, Lapinsky SE, et al. Evaluation of a ventilation strategy to prevent barotrauma in patients at high risk for acute respiratory distress syndrome. Pressure- and Volume-Limited Ventilation Strategy Group. N Engl J Med. 1998;338(6):355-61.
9. Brower RG, Shanholtz CB, Fessler HE, Shade DM, White P Jr, Wiener CM, et al. Prospective, randomized, controlled clinical trial comparing traditional versus reduced tidal volume ventilation in acute respiratory distress syndrome patients. Crit Care Med. 1999;27(8):1492-8.
10. Brower RG, Lanken PN, MacIntyre N, Matthay MA, Morris A, Ancukiewicz M, Schoenfeld D, Thompson BT; National Heart, Lung, and Blood Institute ARDS Clinical Trials Network. Higher versus lower positive end-expiratory pressures in patients with the acute respiratory distress syndrome. N Engl J Med. 2004;351(4):327-36.
11. Meade MO, Cook DJ, Guyatt GH, Slutsky AS, Arabi YM, Cooper DJ, Davies AR, Hand LE, Zhou Q, Thabane L, Austin P, Lapinsky S, Baxter A, Russell J, Skrobik Y, Ronco JJ, Stewart TE; Lung Open Ventilation Study Investigators. Ventilation strategy using low tidal volumes, recruitment maneuvers, and high positive end-expiratory pressure for acute lung injury and acute respiratory distress syndrome: a randomized controlled trial. JAMA. 2008;299(6):837-45.
12. Mercat A, Richmond JC, Vieille B, Jaber S, Osman D, Diehl JL, Lemfrat JY, Prat G, Richette J, Nieszowska A, Gervais C, Baudot J, Bouadma L, Brochard L; Expiratory Pressure (Express) Study Group. Positive end-expiratory pressure setting in adults with acute lung injury and acute respiratory distress syndrome: a randomized controlled trial. JAMA. 2008;299(6):646-55.
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13. Briel M, Meade M, Mercat A, Brower RG, Talmor D, Walter SD, et al. Higher vs lower positive end-expiratory pressure in patients with acute lung injury and acute respiratory distress syndrome: systematic review and meta-analysis. JAMA. 2010;303(9):665-73.

14. Gattinoni L, Caironi P. Refining ventilatory treatment for acute lung injury and acute respiratory distress syndrome. JAMA. 2008;299(6):691-3.

15. Kacmarek RM, Villar J, Sulemanji D, Montiel R, Fernandez C, Blasco J, Koh Y, Soler JA, Martinez D, Hernandez M, Tucci M, Borges JB, Lubillo S, Santos A, Araujo JB, Amato MB, Suarez-Sipmann F. Open Lung Approach Network. Open Lung Approach for the Acute Respiratory Distress Syndrome: A Pilot. Randomized Controlled Trial. Crit Care Med. 2016;44(1):32-42.

16. Protti A, Andreis DT, Monti M, Santini A, Sparacino CC, Langer T, et al. Lung stress and strain during mechanical ventilation: any difference between statics and dynamics? Crit Care Med. 2013;41(4):1046-55.

17. Amato MB, Meade MO, Slutsky AS, Brochard L, Costa EL, Schoenfeld DA, et al. Driving pressure and survival in the acute respiratory distress syndrome. N Engl J Med. 2015;372(8):747-55.

18. ART Investigators. Rationale, study design, and analysis plan of the Alveolar Recruitment for ARDS Trial (ART): study protocol for a randomized controlled trial. Trials. 2012;13:153.

19. Marini JJ. Should We Embrace the “Open Lung” Approach? Crit Care Med. 2013;41(5):1696-7.

20. Gattinoni L, Meade M, Chiumello D, Ranieri VM, Quintel M, et al. Lung recruitment in patients with the acute respiratory distress syndrome. N Engl J Med. 2006;354(17):1775-86.

21. Gattinoni L, Meade M, Santini A, Barbieri M, et al. Lung inhomogeneity in patients with acute respiratory distress syndrome. Am J Respir Crit Care Med. 2014;190(2):149-58.

22. Dreyfuss D, Ricard JD, Gaudry S. Did studies on HFOV fail to improve ARDS mortality? A trial of intraoperative low-tidal-volume ventilation in abdominal surgery. Anesthesiology. 2013;118(6):1307-21.

23. Lavergne P, Selmo G, Lanza C, Chiesa A, Frigerio A, Bacuzzi A, et al. Protective mechanical ventilation during general anaesthesia for open abdominal surgery improves postoperative pulmonary function. Anesthesiology. 2013;118(6):1307-21.

24. PROVE Network Investigators for the Clinical Trial Network of the European Society of Anaesthesiologists, Hemmes SN, Gama de Abreu M, Pelosi P, Schultz MJ. High versus low positive end-expiratory pressure during general anaesthesia for open abdominal surgery (PROVHLD trial): a multicentre randomised controlled trial. Lancet. 2014;384(9942):495-503.

25. Futier E, Constantin JM, Pascal J, Eurin M, Neuschwander-Celis E, Goutte C, Baboi L, Ayzac L; PROSEVA Study Investigators. Neuromuscular blockers in early acute respiratory distress syndrome. N Engl J Med. 2010;363(12):1107-16.

26. Retamal J, Bergamini BC, Carvalho AR, Bozza FA, Borzone G, Borges JB, et al. Non-lobar atelectasis generates inflammation and structural alveolar injury in the surrounding healthy tissue during mechanical ventilation. Crit Care. 2014;18(5):505.

27. Retamal J, Bugedo G, Diquelme F, Varas J, Retamal J, Besa C, et al. Tidal volume is a major determinant of cyclic recruitment-de-recruitment in acute respiratory distress syndrome. Minerva Anestesiol. 2011;77(4):418-26.

28. Bein T, Weber-Carstens S, Goldmann A, Müller T, Staudinger T, Brederlau J, et al. Lower tidal volume strategy (approximately 3 ml/kg) combined with extracorporeal CO2 removal versus “conventional” protective ventilation (6 ml/kg) in severe ARDS: the prospective randomized Xtravent-study. Intensive Care Med. 2013;39(5):847-56.

29. Retamal J, Libuy J, Jiménez M, Delgado M, Besa C, Bugedo G, et al. Preliminary study of ventilation with 4 ml/kg tidal volume in acute respiratory distress syndrome: feasibility and effects on cyclic recruitment - derecruitment and hyperinflation. Crit Care. 2013;17(1):R16.

30. Papazian L, Forel JM, Gacouin A, Penot-Ragon C, Perrin G, Loundou A, Jaber S, Arnal JM, Perez D, Seghbyoun JM, Constantin JM, Courant P, Lefrant JY, Guerin C, Prat G, Morange S, Bengler C, Richecoeur J, Rosselli D, Ayzac L; PROSEVA Study Investigators. Neurromuscular blockers in early acute respiratory distress syndrome. Rev Bras Ter Intensiva. 2012;24(1):43-51.

31. Bugedo G, Bruhn A, Regueira T, Romero C, Retamal J, Hernández G. Positive end-expiratory pressure increases strain in patients with ALI/ARDS. Rev Bras Ter Intensiva. 2012;24(1):43-51.

32. Retamal J, Bugedo G, Larsson A, Bruhn A. High PEEP levels are associated with overdistension and tidal recruitment/derecruitment in ARDS patients. Acta Anaesthesiol Scand. 2015;59(9):1161-9.

33. Young D, Lam SE, Shah S, MacKenzie I, Tunnicliffe W, Lall R, Rowan K, Cuthbertson BH, OSCILLATE Study Group. High-frequency oscillation for acute respiratory distress syndrome. N Engl J Med. 2013;368(9):806-13.

34. Ferguson ND, Cook DJ, Guyatt GH, Mehta S, Hand L, Austin P, Zhou Q, Matte A, Walter SD, Lamontagne F, Granott JT, Arabi YM, Arroliga AC, Stewart TE, Slutsky AS, Meade MO. OSCILLATE Trial Investigators; Canadian Critical Care Trials Group. High-frequency oscillation in early acute respiratory distress syndrome. N Engl J Med. 2013;368(9):795-805.

35. Determann RM, Royakkers A, Wothhua EK, Vlaar AP, Choi G, Paulus F, et al. Lung stress and strain during mechanical ventilation: any difference between statics and dynamics? Crit Care Med. 2013;41(4):1046-55.

36. Bruhn A, Bugedo G, Diquelme F, Varas J, Retamal J, Besa C, et al. Tidal volume is a major determinant of cyclic recruitment-de-recruitment in acute respiratory distress syndrome. Minerva Anestesiol. 2011;77(4):418-26.

37. Terragni PP, Del Sorbo L, Mascia L, Urbino R, Martin EL, Birocco A, et al. Tidal volume lower than 6ml/kg enhances lung protection: role of extracorporeal carbon dioxide removal. Anesthesiology. 2009;111(4):826-35.

38. Bein T, Weber-Carstens S, Goldmann A, Müller T, Staudinger T, Brederlau J, et al. Lower tidal volume strategy (approximately ≈3 ml/kg) combined with extracorporeal CO2 removal versus “conventional” protective ventilation (6 ml/kg) in severe ARDS: the prospective randomized Xtravent-study. Intensive Care Med. 2013;39(5):847-56.

39. Retamal J, Libuy J, Jiménez M, Delgado M, Besa C, Bugedo G, et al. Preliminary study of ventilation with 4 ml/kg tidal volume in acute respiratory distress syndrome: feasibility and effects on cyclic recruitment - derecruitment and hyperinflation. Crit Care. 2013;17(1):R16.

40. Papazian L, Forel JM, Gacouin A, Penot-Ragon C, Perrin G, Loundou A, Jaber S, Arnal JM, Perez D, Seghbyoun JM, Constantin JM, Courant P, Lefrant JY, Guerin C, Prat G, Morange S, Bengler C, Richecoeur J, Rosselli D, Ayzac L; PROSEVA Study Investigators. Neurromuscular blockers in early acute respiratory distress syndrome. Rev Bras Ter Intensiva. 2012;24(1):43-51.