High positive end-expiratory pressure: only a dam against oedema formation?

Alessandro Protti1*, Davide T Andreis2, Giacomo E Iapichino2, Massimo Monti2, Beatrice Comini2, Marta Milesi2, Loredana Zani2, Stefano Gatti3, Luciano Lombardi4 and Luciano Gattinoni1,2

Abstract

Introduction: Healthy piglets ventilated with no positive end-expiratory pressure (PEEP) and with tidal volume (VT) close to inspiratory capacity (IC) develop fatal pulmonary oedema within 36 h. In contrast, those ventilated with high PEEP and low VT, resulting in the same volume of gas inflated (close to IC), do not. If the real threat to the blood-gas barrier is lung overinflation, then a similar damage will occur with the two settings. If PEEP only hydrostatically counteracts fluid filtration, then its removal will lead to oedema formation, thus revealing the deleterious effects of overinflation.

Methods: Following baseline lung computed tomography (CT), five healthy piglets were ventilated with high PEEP (volume of gas around 75% of IC) and low VT (25% of IC) for 36 h. PEEP was then suddenly zeroed and low VT was maintained for 18 h. Oedema was diagnosed if final lung weight (measured on a balance following autopsy) exceeded the initial one (CT).

Results: Animals were ventilated with PEEP 18 ± 1 cmH2O (volume of gas 875 ± 178 ml, 89 ± 7% of IC) and VT 213 ± 10 ml (22 ± 5% of IC) for the first 36 h, and with no PEEP and VT 213 ± 10 ml for the last 18 h. On average, final lung weight was not higher, and actually it was even lower, than the initial one (284 ± 62 vs. 347 ± 36 g; P = 0.01).

Conclusions: High PEEP (and low VT) do not merely impede fluid extravasation but rather preserve the integrity of the blood-gas barrier in healthy lungs.

Introduction

Mechanical ventilation is a pivotal therapy for respiratory failure, although overinflation may injure the lung [1-3].

We have recently shown that healthy piglets ventilated with no positive end-expiratory pressure (PEEP) and with tidal volume (VT) close to inspiratory capacity (IC) die with inflammatory pulmonary oedema within 36 h. In contrast, those ventilated with high PEEP (≥18 cmH2O; volume of gas around 75% of IC) and low VT (25% of IC) survive with normal lungs for 54 h [4].

Inflammatory pulmonary oedema develops when lung capillary transmural (internal minus external) pressure drives excessive fluid filtration through a disrupted, highly permeable, blood-gas barrier [5]. As a result, exudates accumulate in the extravascular space and lung weight increases [6,7]. If overinflation per se is the real threat to the blood-gas barrier [8], then ventilation with large VT alone (100% of IC) will be as harmful as ventilation with high PEEP (75% of IC) and low VT (25% of IC). In fact, in both cases, the volume of gas globally inflated will be equal to IC. However, high PEEP (and low VT) will possibly impede extravasation by lowering venous return, cardiac output and pulmonary capillary inflow (and pressure) while increasing extravascular pressure [9,10]. Oedema will then not develop even if the blood-gas barrier loses its integrity.

To test this hypothesis, we ventilated healthy piglets with high PEEP (and low VT) (as above) and then suddenly removed PEEP to allow free extravasation through a possibly disrupted blood-gas barrier. Lung weight was expected to increase as oedema formation was no longer impeded.
Materials and methods
The study complied with international recommendations [11] and was approved by the Italian Ministry of Health (protocol number 01/10).

Five healthy, sedated and paralysed piglets (21 ± 1 kg) were surgically prepared in supine position. They were then turned prone and ventilated with no PEEP and with a VT of 10 ml/kg of body weight (Engström Carestation, GE Healthcare; Madison, WI, USA). Lung computed tomography (CT) was taken at 0 cmH2O (functional residual capacity, FRC), around 18 cmH2O (the level of PEEP planned to be used) (see below) and 45 cmH2O (arbitrarily defined as total lung capacity, TLC) of airway pressure. Lung gas volumes and weight were measured by quantitative analysis [12].

Following this pre-study period, PEEP was set around 18 cmH2O while VT was kept constant (PEEP phase). Based on a previous study [4], we expected that the volume of gas so inflated in the form of PEEP would have been around 75% and VT around 25% of IC (TLC minus FRC). Therefore, the volume of gas inflated by end of inspiration (volume of gas due to PEEP plus VT) was predicted to be equal to IC and end-inspiratory lung volume, including FRC, to approach TLC. This setting was kept constant for 36 h (PEEP phase). PEEP was then suddenly zeroed and VT was maintained for 18 h (ZEEP phase) (Figure 1).

Results of the lung CT scan analysis became available once the study had already started. They did not influence the setting described above but permitted exact computation of volumes of gas effectively inflated (as PEEP and VT) in relation to lung capacities. During the entire 54-h study period the animals were kept prone, in 10° Trendelenburg position [13]. Respiratory rate was 15 breaths per minute, inspiratory-to-expiratory time ratio 1:2 (1:3 if intrinsic PEEP developed) and inspired oxygen fraction was 0.50. Tracheal suctioning and recruitment manoeuvres were only performed before applying PEEP and after its removal. Use of normal saline and norepinephrine was standardised to try to maintain mean arterial pressure above 60 mmHg. Respiratory system and lung mechanics, blood gas analysis and haemodynamics (including cardiac output by thermodilution) were assessed every 6 h. Data were also collected during the pre-study period (baseline) and soon after PEEP removal (time 0 of the ZEEP phase). Transpulmonary pressure was computed as (end-inspiratory airway pressure) - [(end-inspiratory oesophageal pressure) - (oesophageal pressure at 0 cmH2O of airway pressure)]. Transmural pulmonary arterial and pulmonary artery occlusion pressures were computed as (end-expiratory intravascular pressures) - [(end-expiratory oesophageal pressure) - (oesophageal pressure at 0 cmH2O of airway pressure)] [14]. Urinary output was recorded every hour and water balance was computed as the difference between saline infusion and urinary output.

At the end of the study, animals were sacrificed (potassium chloride, 40 mEq intravenous) and exsanguinated. Lungs were excised and weighted on a balance. This (final) lung weight was compared with the initial one, measured on pre-study CT taken at 0 cmH2O of airway pressure (the degree of agreement between these two methods for measuring lung weight was assessed in five other piglets, as reported in Additional file 1). Pulmonary oedema was diagnosed if lung weight had increased across the entire study period. The right lung was used for the calculation of wet-to-dry weight ratio and blind histological examination [15]. Twelve piglets uneventfully ventilated for 54 h with no PEEP and low VT (as part of a previous study [3]) that did not develop pulmonary oedema were used as controls.

Statistical analysis
Numerical data are reported as mean ± standard deviation. Based on distribution (Shapiro-Wilk test), they were analysed using Student’s (paired) t test or Mann-Whitney rank sum test, one-way repeated measures analysis of variance (RM ANOVA) or RM ANOVA on ranks. Post hoc comparisons were done using the Holm-Sidak or Dunn’s method. Categorical data are reported as median (interquartile range) and were analysed using the Mann-Whitney rank sum test. A P value <0.05 was considered significant (SigmaPlot 11.0, Jandel Scientific Software; San Jose, CA, USA).

For more information on methods, please refer to Additional file 2.

Results
Animals were ventilated with PEEP 18 ± 1 cmH2O and VT 213 ± 10 ml for the first 36 h (PEEP phase) and with no PEEP and VT 213 ± 10 ml for the last 18 h (ZEEP phase).

On baseline lung CT (Additional file 3), application of PEEP 18 ± 1 cmH2O resulted in inflation of 875 ± 178 ml of gas, equal to 89 ± 7% of estimated IC (988 ± 176 ml). VT was 22 ± 5% of estimated IC. The volume of gas inflated by end of inspiration during the PEEP phase (volume of gas due to PEEP plus VT) was slightly higher than estimated IC (1088 ± 177 vs. 988 ± 176 ml, P = 0.05). End-inspiratory lung volume, including FRC (425 ± 66 ml), was slightly higher than estimated TLC (1512 ± 229 vs. 1413 ± 225 ml, P = 0.05).

Changes in systemic and pulmonary haemodynamics are shown in Figure 2. Mean arterial pressure (and cardiac output) diminished soon after the application of PEEP despite aggressive fluid resuscitation and norepinephrine infusion (Additional file 4), but returned to baseline before...
PEEP was removed. Transmural mean pulmonary arterial pressure did not change over time while transmural pulmonary artery occlusion pressure tended to be higher than pre-study values following PEEP removal. By the end of the experiment, water balance ranged between +1600 and +6400 ml and body weight had accordingly increased (from 21 ± 1 to 25 ± 2 kg, \(P = 0.01\)).

Respiratory system mechanics, lung mechanics and gas exchange did not deteriorate during the PEEP phase (Additional file 5). Nonetheless, data recorded immediately after PEEP removal were slightly worse than those collected at baseline; as far as subsequent oedema formation was concerned, no further change occurred during the ZEEP phase (Table 1).

At autopsy, lungs looked pink and normally inflated, except for small areas of atelectasis (Additional file 6). Final lung weights did not exceed initial ones (Figure 3). Lung wet-to-dry weight ratio (5.1 ± 0.3 vs. 4.8 ± 0.7, \(P = 0.38\)) and histology (Table 2) did not differ from those of controls.

**Discussion**

Healthy piglets ventilated with high PEEP and low \(V_T\) for 36 h do not develop pulmonary oedema when airway pressure is suddenly lowered (by removing PEEP). This proves that high PEEP and low \(V_T\) do not permanently alter the integrity of the alveolar-capillary interface even if lung inflation approaches its upper physiological limit (volume of gas globally inflated equal to IC).

We have previously shown that healthy piglets ventilated for 54 h with high PEEP (never removed) and low \(V_T\) survive with normal lungs [4]. There, lack of pulmonary oedema could have been attributed to diminished transmural pulmonary capillary pressure (that drives fluid extravasation) across an abnormally highly permeable (overstretched) blood-gas barrier. In fact, application of high PEEP always caused hypotension and low cardiac output while lungs were maximally inflated [4].

PEEP may or may not prevent lung oedema (or even worsen it) depending on relative changes in pulmonary intravascular and extravascular (alveolar and extra-alveolar)

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**Figure 1 Study design.** Initial lung weights were measured by CT. Animals were then ventilated with high PEEP (volume of gas around 75% of IC; dotted line) and low VT (25% of IC; spikes) for 36 h (PEEP phase). Thereafter, PEEP was suddenly zeroed and low VT maintained for 18 additional h (ZEEP phase). Animals were then sacrificed and final lung weights were measured on a balance. B, baseline; CT, computed tomography; FRC, functional residual capacity; IC, inspiratory capacity; PEEP, positive end-expiratory pressure; TLC, total lung capacity; \(V_T\), tidal volume; ZEEP, zero end-expiratory pressure.
pressures, lung surface area (through which filtration occurs), pulmonary endothelium and epithelium porosity [16-20]. Since its initial use, PEEP was mainly thought to tamponade great veins (thus lowering pulmonary blood flow and pressure) and ‘exert an opposing force tending to hinder the outpouring of both red corpuscles and serum from the pulmonary capillaries’ [21]. Other means of lowering intravascular pulmonary pressure, such as phlebotomy or vasodilatation, can attenuate pulmonary oedema formation, even if the blood-gas barrier is abnormally

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**Figure 2** Haemodynamics. Mean arterial pressure (panel A, n = 5), cardiac output (panel B, n = 5), transmural pulmonary arterial pressure (panel C, n = 4) and transmural pulmonary artery occlusion pressure (panel D, n = 4) were recorded at baseline (B), during ventilation with high PEEP and low VT (36 h) and during ventilation with no PEEP (ZEEP) and low VT (18 h). During ventilation with high PEEP, oesophageal pressure at 0 cmH2O was assumed to have changed linearly from the value recorded at B (at 0 cmH2O) to the value recorded at time 0 of the ZEEP phase. *P* values refer to one-way RM ANOVA (on ranks if appropriate). *P* <0.05 vs. B (Holm-Sidak or Dunn’s method). B, baseline; PEEP, positive end-expiratory pressure; RM ANOVA, repeated measures analysis of variance; VT, tidal volume; ZEEP, zero end-expiratory pressure.

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**Table 1 Lung function**

|                     | Baseline       | Ventilation with no PEEP (ZEEP phase) | 0 h   | 6 h   | 12 h  | 18 h  |  P    |
|---------------------|----------------|--------------------------------------|-------|-------|-------|-------|-------|
| **End-inspiratory airway pressure (cmH2O)** | 10 ± 1         | 14 ± 3*                               | 13 ± 2* | 13 ± 3* | 13 ± 3* | 0.002 |
| **Transpulmonary pressure (cmH2O)**          | 5 ± 2          | 10 ± 3*                               | 9 ± 3*  | 9 ± 3*  | 10 ± 4* | 0.001 |
| **Arterial oxygen tension (mmHg)**           | 222 ± 43       | 187 ± 58                              | 181 ± 44 | 180 ± 47 | 186 ± 24 | 0.11  |
| **Arterial carbon dioxide tension (mmHg)**   | 41 ± 3         | 36 ± 4                                | 39 ± 4  | 38 ± 3  | 39 ± 3  | 0.16  |

Respiratory system mechanics, lung mechanics and gas exchange at baseline and during ventilation with no PEEP (ZEEP phase) (18 h). End-inspiratory airway pressure was recorded during a 5-sec end-inspiratory pause. *P* values refer to one-way RM ANOVA (on ranks if appropriate). *P* <0.05 vs. baseline (Holm-Sidak or Dunn’s method). PEEP, positive end-expiratory pressure; ZEEP, zero end-expiratory pressure; RM ANOVA, repeated measures analysis of variance.
permeable [22,23]. On the other side, increasing transmural pulmonary capillary pressure with dopamine [8] or fluids [24-26] aggravates inflammatory pulmonary oedema and sudden removal of a previously existing backward pressure on the pulmonary capillaries (that is PEEP) is followed by an increased permeability of the capillary wall and leakage of serum [21]. Therefore, lack of oedema per se does not prove that high PEEP and low VT are safe.

In this present work, we specifically addressed one of the mechanisms possibly underlying our [4] and others’ [6,8] previous findings (namely, diminished pulmonary capillary filtration pressure) by abruptly removing a potential obstacle to oedema formation (high PEEP itself). We ventilated healthy piglets with high PEEP and low VT (volume of gas globally inflated around estimated IC) for 36 h. By this time, animals ventilated with large VT alone (always resulting in the same end-inspiratory lung volume) develop overt pulmonary oedema, as indicated by heavy and congested lungs, altered respiratory system mechanics and gas exchange, and pulmonary and systemic inflammation [3]. Since lung injury may not become macroscopically evident as long as PEEP counteracts fluid filtration, PEEP was then suddenly zeroed. Fluid filtration should have freely occurred, driven by normal (and occasionally supra-normal) haemodynamics, had the blood-gas barrier been disrupted [18,24,27].

Acting similarly, we have previously demonstrated a small, non-significant increase in lung weight following PEEP removal in four healthy piglets, finally ventilated with low VT for 3 h [4]. In order to exclude that overt pulmonary oedema could have developed had the experiments lasted longer, we decided to extend the duration of ventilation with low VT and no PEEP to 18 h. Even so, lung weight did not increase (and it actually decreased, possibly as a consequence of exsanguination [28]) and mechanics and gas exchange did not deteriorate with time. This proves that ventilation with high PEEP and low VT over the previous 36 h had not grossly and permanently altered the permeability of the blood-gas barrier. The minor decrease in pulmonary compliance and arterial oxygenation that became evident when PEEP was removed (and did not worsen over time) was possibly related to

![Figure 3 Lung weight](http://ccforum.com/content/17/4/R131)

Figure 3 Lung weight. Individual changes in lung weight across the entire study period. Straight lines are used to connect individual points (not to suggest that changes occurred linearly over time). Initial lung weights were measured by quantitative analysis of lung CT obtained at baseline; final lung weights were measured on a balance following autopsy. *P = 0.01 (Student’s paired t test). CT, computed tomography.
VT (and high inspiratory flows) may have a major role as inflation, are not known. On one side, avoidance of large pulmonary oedema, despite resulting in extremely large lung proliferations.

Alveolar type II pneumocytes proliferation 3
Alveolar macrophage proliferation 2
Alveolar neutrophil infiltration 2
Alveolar haemorrhage 1
Interstitial congestion 2
Emphysematous changes 2

Animals ventilated with no PEEP and low VT for 54 h (present study) were compared with those of animals ventilated with no PEEP and low VT for 48 h (previous study) [3]. Alterations were graded from 0 to 4, with higher scores indicating more severe and diffuse abnormalities. Total score was the sum of all individual scores. P values refer to Mann-Whitney rank sum test. PEEP, positive end-expiratory pressure; VT, tidal volume.

Results of animals ventilated with high PEEP and low VT for 36 h and with no PEEP and low VT for 18 h (present study) were compared with those of animals ventilated with no PEEP and low VT for 54 h (previous study) [3]. Alveolar type II pneumocytes proliferation 2
Alveolar neutrophil infiltration 2
Alveolar haemorrhage 1
Interstitial fibrosis 1
Interstitial lymphocytes proliferation 2
Interstitial thickening 2
Hyline membranes formation 0
Organization of alveolar exudate 1
Total score 18

Key messages
• Healthy lungs ventilated with high PEEP and low VT do not develop pulmonary oedema, even if globally inflated up to their total capacity.
• High PEEP (and low VT) do not merely hydrostatically counteract pulmonary fluid extravasation.
• Mechanical ventilation with high PEEP and low VT does not grossly injure the blood-gas barrier in healthy lungs.

Additional material

Additional file 1: Bland-Altman plot of lung weight measured with balance vs. CT scan. Five other healthy piglets (18 ± 1 kg) underwent lung CT at 0 cmH2O of airway pressure and were then sacrificed and exsanguinated. Results of quantitative analysis of CT scan were compared with excised lung weight, measured on a balance. The Bland-Altman plot of lung weight measured with the two techniques show a bias -91 g.
Additional file 2: Additional methods. Further details on surgical preparation, quantitative analysis of lung CT, haemodynamic protocol, sacrifice and autopsy are shown. CT, computed tomography.

Additional file 3: Baseline lung CT. Results of quantitative analysis of lung CT scans, which allows for exact computation of lung volumes and capacity of gas due to PEEP. CT, computed tomography; PEEP, positive end-expiratory pressure.

Additional file 4: Additional haemodynamic variables throughout the experiment. Urinary output (panel A, n = 5), water balance (panel B, n = 5), urinary electrolytes (panel C and D, n = 4), arteriovenous oxygen difference (panel E, n = 5), blood lactate (panel F, n = 5), and rate of norepinephrine infusion (panel G, n = 5) were recorded at baseline (B), during ventilation with high PEEP and low Vt (36 h) and during ventilation with no PEEP (ZEEP) and low Vt (18 h). P values refer to one-way RM ANOVA (on ranks if appropriate); *P < 0.05 vs. B (Holm-Sidak or Dunn’s method). B, baseline; PEEP, positive end-expiratory pressure; RM ANOVA, repeated measures analysis of variance; Vt, tidal volume; ZEEP, zero end-expiratory pressure.

Additional file 5: Lung function during ventilation with high PEEP and low Vt. Respiratory system mechanics (panel A) lung mechanics (panel B), and gas exchange (panel C and D) were recorded during 36 h of ventilation with high PEEP and low Vt. Dorsophaegal pressure at 0 cmH2O of airway pressure was assumed to have changed linearly from the value recorded at baseline (B) (at 0 cmH2O) to the value recorded at time 0 of the ZEEP phase. P values refer to one-way RM ANOVA (on ranks if appropriate); *P < 0.05 vs. B (Holm-Sidak or Dunn’s method). B, baseline; PEEP, positive end-expiratory pressure; RM ANOVA, repeated measures analysis of variance; Vt, tidal volume, ZEEP, zero end-expiratory pressure.

Additional file 6: Autopotic lung appearance. Lungs of animals ventilated with high PEEP and low Vt for 36 h and with no PEEP and low Vt (18 h) (experimental study) are shown in panel A. For comparison, lungs of animals ventilated for 54 h with no PEEP and low Vt (lung weight changed from 377 to 220 g) (panel B), with no PEEP and large Vt (close to inspiratory capacity) (lung weight increased from 395 to 721 g) (panel C) and with high PEEP and low Vt for 54 h (lung weight changed from 282 to 290 g) (panel D) are shown. PEEP, positive end-expiratory pressure; Vt, tidal volume.

Abbreviations
CT: computed tomography; FRC: functional residual capacity; IC: inspiratory capacity; PEEP: positive end-expiratory pressure; RM ANOVA: repeated measures analysis of variance; TLC: total lung capacity; Vt: tidal volume; ZEEP: zero end-expiratory pressure.

Competing interests
This study was funded in part by GE Healthcare. The authors declare that they have no competing interests that can have influenced the submitted manuscript.

Authors’ contributions
AP participated in the study design, statistical analysis and wrote the manuscript. DTA participated in the study design, ran the experiments, performed statistical analysis and revised the manuscript. GEI participated in the study design, ran the experiments, performed statistical analysis and revised the manuscript. MM participated in the study design and data acquisition. BC participated in data acquisition and analysis. MM participated in data acquisition and analysis. LZ participated in data acquisition and analysis. SG performed the surgical preparation and revised the manuscript. LL performed the lung computed tomography and participated in the study design. LG participated in the study design and revised the manuscript. All authors read and approved the final version of the manuscript.

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Authors’ details
1Dipartimento di Anestesia, Rianimazione (Intensiva e Subintensiva) e Terapia del Dolore, Fondazione IRCCS Ca’ Granda - Ospedale Maggiore Policlinico, via Francesco Sforza, 35, 20122 Milan, Italy. 2Dipartimento di Fisiopatologia Medico-Chirurgica e dei Trapianti, Università degli Studi di Milano, Via Francesco Sforza, 35, 20122 Milan, Italy. 3Centro di Ricerche Chirurgiche Precliniche, Fondazione IRCCS Ca’ Granda - Ospedale Maggiore Policlinico, Università degli Studi di Milano, Via Francesco Sforza, 35, 20122 Milan Italy. 4Dipartimento di Radiologia, Fondazione IRCCS Ca’ Granda - Ospedale Maggiore Policlinico, via Francesco Sforza, 35, 20122 Milan, Italy.

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