Flow-controlled ventilation during EVLP improves oxygenation and preserves alveolar recruitment

Sofie Ordies\textsuperscript{1,2,3*}, Michaela Orlitova\textsuperscript{1,3,4}, Tobias Heigl\textsuperscript{3,5}, Annelore Sacreas\textsuperscript{3,5}, Anke Van Herck\textsuperscript{3,5,6}, Janne Kaes\textsuperscript{3,5}, Berta Saez\textsuperscript{3,5,6}, Arno Vanstapel\textsuperscript{3,5,7}, Laurens Ceulemans\textsuperscript{3,4,5}, Bart M. Vanaudenaerde\textsuperscript{3,5}, Robin Vos\textsuperscript{3,5,6}, Johny Verschakelen\textsuperscript{8}, Geert M. Verleden\textsuperscript{3,5,6}, Stijn E. Verleden\textsuperscript{3,5}, Dirk E. Van Raemdonck\textsuperscript{3,4,5} and Arne P. Neyrinck\textsuperscript{1,2,3*}

\textbf{Background}

Lung transplantation (LTx) remains the last-resort solution for patients suffering from end-stage respiratory diseases \cite{1}. However, the lack of suitable organs remains an important limiting factor for survival of wait listed candidates.

\textbf{Abstract}

\textbf{Background:} Ex vivo lung perfusion (EVLP) is a widespread accepted platform for preservation and evaluation of donor lungs prior to lung transplantation (LTx). Standard lungs are ventilated using volume-controlled ventilation (VCV). We investigated the effects of flow-controlled ventilation (FCV) in a large animal EVLP model.

Fourteen porcine lungs were mounted on EVLP after a warm ischemic interval of 2 h and randomized in two groups (\(n=7\)/group). In VCV, 7 grafts were conventionally ventilated and in FCV, 7 grafts were ventilated by flow-controlled ventilation. EVLP physiologic parameters (compliance, pulmonary vascular resistance and oxygenation) were recorded hourly. After 6 h of EVLP, broncho-alveolar lavage (BAL) was performed and biopsies for wet-to-dry weight (W/D) ratio and histology were taken. The left lung was inflated, frozen in liquid nitrogen vapors and scanned with computed tomography (CT) to assess regional distribution of Hounsfield units (HU).

\textbf{Results:} All lungs endured 6 h of EVLP. Oxygenation was better in FCV compared to VCV (\(p=0.01\)) and the decrease in lung compliance was less in FCV (\(p=0.03\)). W/D ratio, pathology and BAL samples did not differ between both groups (\(p=0.16\), \(p=0.55\) and \(p=0.62\)). Overall, CT densities tended to be less pronounced in FCV (\(p=0.05\)). Distribution of CT densities revealed a higher proportion of well-aerated lung parts in FCV compared to VCV (\(p=0.01\)).

\textbf{Conclusions:} FCV in pulmonary grafts mounted on EVLP is feasible and leads to improved oxygenation and alveolar recruitment. This ventilation strategy might prolong EVLP over time, with less risk for volutrauma and atelectrauma.

\textbf{Keywords:} Lung transplantation, Ex vivo lung perfusion, Flow-controlled ventilation, Donor management, Volume-controlled ventilation, Porcine large animal model

\textbf{Background}

Lung transplantation (LTx) remains the last-resort solution for patients suffering from end-stage respiratory diseases \cite{1}. However, the lack of suitable organs remains an important limiting factor for survival of wait listed candidates.
Ex vivo lung perfusion (EVLP) is a form of machine perfusion and has been introduced in the field of LTx to evaluate graft function outside the body [2]. During EVLP, the lungs are perfused with a normothermic solution and at the same time ventilated with positive pressure. Graft function is mainly monitored using standard physiological variables including compliance, pulmonary vascular resistance (PVR) and oxygenation. EVLP has the potential to assess organs physiologically and biologically prior to transplantation and facilitates longer preservation times compared to conventional hypothermic storage on ice [3]. In the future, active resuscitation of injured grafts by EVLP might be possible, but search of adequate therapeutic strategies is still ongoing [4].

To date, duration of EVLP is still limited up to 12 h clinically [5] and to 24 h in a large animal model [6–8]. A potential reason why EVLP is still limited in time might be the technology itself. The concept of EVLP is based on a physiological environment and assessment. However, the ex situ context cannot be completely compared to in vivo physiology. First, the graft is mounted in a static supine position in the organ chamber. We have demonstrated that prone position alters region distribution of edema accumulation [9]. Second, the graft is mechanically ventilated with positive airway pressure. Patients who receive mechanical ventilation with increased pressures (barotrauma), large volumes (volutrauma), repetitive alveolar opening and closing (atelectrauma) and asynchronous breathing (self-inflicting lung injury) are at risk for developing ventilator-induced lung injury (VILI), which leads to inferior outcome [10–12]. Therefore, one can presume that similar mechanisms of VILI might be important during ex vivo lung perfusion.

The two most frequently used ventilation modes during EVLP are volume-controlled ventilation (VCV) and pressure-controlled ventilation (PCV) [4]. In addition, it is not clear what would be the optimal recruitment maneuver to actively open collapsed alveoli, especially with absence of chest wall. These ventilation strategies only control the inspiratory and not the expiratory phase of the respiratory cycle. Expiration depends on passive elastic recoil of the lungs.

Flow-controlled ventilation (FCV) is an innovative ventilation mode providing a constant flow during both inspiration and expiration, resulting in a linear increase and decrease in intratracheal and thus intrapulmonary pressure. FCV has been shown to enhance lung aeration, to prevent atelectasis and to reduce lung damage [13]. The airway/intratracheal pressure, gas flow and tidal volume patterns of VCV and FCV ventilation modes are illustrated in Fig. 1.

The effects of FCV ventilation have not been previously investigated in a large animal EVLP model. Therefore, the aim of this study was to explore effects of FCV during EVLP compared to conventional VCV. We hypothesized that FCV could improve alveolar recruitment and reduce VILI during EVLP compared to VCV.

Methods

Animals

Fourteen specific pathogen-free domestic male pigs (Topig20, Tojapigs, Escharen, The Netherlands) (median weight 39 kg (36.5–39.6 IQR), n = 7/group) were included in this study after approval by the Ethics Committee (P145/2016). Each animal received human care conform the “Principles of Laboratory Animal Care”, formulated by the National
Society for Medical Research and “Guide for the Care and Use of Laboratory Animals,” prepared by the Institute of Laboratory Animal Resources and reported by the National Institutes of Health, USA (NIH Publication No. 86-23, revised 1996).

**Injury model and EVLP setup**

Animals were sedated and anaesthetized as previously described [9]. Baseline venous and arterial blood samples were taken at 100% fraction of inspiratory oxygen (FiO₂). Baseline respiratory and hemodynamic parameters were recorded with ICM⁺® (Cambridge Enterprise, Cambridge, UK). Donor animals were ventilated with volume-controlled ventilation (VCV) 8 mL/kg, an inspiratory time-to-expiratory ratio (I:E) of 1:2 and a positive end-expiratory pressure (PEEP) of 5 cm H₂O, and respiratory rate (RR) was dependent on maintenance of expiratory carbon dioxide (CO₂) levels between 40 and 45 mmHg. Median sternotomy was performed and heparin (300 U/kg) (LEO Pharma BV, Amsterdam, The Netherlands) was administered prior to introduction of cardiac fibrillation (10 V until witnessed cardiac arrest).

After cardiac arrest, lungs were left untouched and deflated in situ for 2 h of warm ischemia. Sequentially, the grafts were retrogradely flushed with 1.5 L OCS™ solution (Transmedics, Andover, MA, USA). The first 0.5 L flush solution was administered at room temperature (21 °C), the next 1 L was at 4 °C. The lungs were inflated, stored on ice and prepared for EVLP. Our EVLP setup was as follows: acellular perfusate, closed

![Diagram](image-url)  
*Fig. 1: Volume-controlled ventilation (VCV) and flow-controlled ventilation (FCV). Visualization of the airway/tracheal pressures, flow and volume during conventional volume-controlled ventilation (VCV, left) and flow-controlled ventilation (FCV, right) during inspiration and expiration.*
atrium and flow calculated as 40% of cardiac output. A centrifugal pump, heat exchanger, gas exchanger, and reservoir enabled normothermic (37 °C), double-lung EVLP during 6 h as previously reported [9]. Briefly, the lungs were cannulated with XVIVO™ Lung Cannula Set (XVIVO Perfusion™, Göteborg, Sweden).

Study groups and ventilation strategy
Lungs were randomized in two groups (n=7/group). In the VCV group, donor lungs were ventilated according to standard VCV during EVLP (tidal volume (TV) of 7 mL/kg; I:E 1:2, 5 cm H2O PEEP and RR of 7 breaths/min) (Aestiva 3000; GE Healthcare Europe GmbH, Little Chalfont, UK) and an endotracheal tube (ETT) of 8.0 mm. A recruitment maneuver based on an increase of PEEP to 10 cm H2O and inspiratory pressure to 25 cm H2O during 60 s was applied.

In the FCV group, grafts were ventilated with FCV (Evone®, Ventinova Medical, Eindhoven, The Netherlands) during EVLP. The settings of FCV are based on an adjustment of the gas flow, PEEP and inspiratory pressure only. Ventilatory rate and volume cannot be independently set and are dependent on the driving pressure (inspiratory pressure–PEEP) and flow. For these experiments, the driving pressure was set to reach a TV of 7 mL/kg and the flow adjusted to reach a RR of 7 breaths/min. I:E was 1:1 and 5 cm H2O PEEP was maintained during 6 h of EVLP. A special requirement to control the gas flow, is the use of a narrow-bore ETT with direct measurement of intratracheal pressure (Tritube®; Ventinova Medical, Eindhoven, The Netherlands). In both groups, FiO2 during EVLP was 21%. Hourly blood gases were taken with a FiO2 of 100%.

Evaluation of graft function and lung injury
All lungs remained on EVLP for 6 h. During EVLP, hemodynamic [pulmonary artery pressure (PAP), left atrial pressure (LAP)] and respiratory parameters (compliance, RR, flow) were monitored. Since compliance was calculated differently, based on extra-tracheal airway pressure proximally of the ETT in VCV and intratracheal pressure in FCV, static compliance was recalculated in the VCV group using following formula expiratory TV/(P_{plateau}−PEEP). For FCV, dynamic compliance calculated by Evone® was based on inspiratory tidal volume measured by mass flowmetry and driving pressure. Delta (Δ) of PVR and oxygenation and compliance were calculated as difference of the respective parameter between 6 and 1 h of EVLP.

At the end of EVLP, a ventral and dorsal biopsy of the right lung were taken for pathology and for calculation of wet-to-dry weight (W/D) ratio. Biopsies were processed as previously reported [9], hematoxylin and eosin staining was scored by a pathologist blinded for group information. A composite score for pathology was used, including congestion, interstitial thickening and alveolar edema [each parameter scored between 0 (absent) and 3 (manifest present)]. The presence of neutrophils was noted as 0 (absent) or 1 (present). The sum of all scores was calculated and divided by the maximum score 20 to illustrate the percentage of injury. Lungs were weighed prior to and after EVLP. Samples for W/D were mounted in a heated oven of 80 °C and weighed after 72 h. After EVLP, a broncho-alveolar lavage (BAL) sample was taken in the right middle lobe by applying two times 30 mL of saline 0.9%. Pooled fractions were obtained and a 100µL cytospin was stained with Diff-Quick (Dade Behring, Newark, NJ, USA) to calculate
total and differential cell counts [14]. Finally, the left lung was inflated at a constant inspiratory pressure of 25 cm H$_2$O and frozen into liquid nitrogen fumes and computed tomography (CT) scanned (Siemens Somatom scanner, Erlangen, Germany) at 120 kV and 110 mAs. HOROS® [DICOM Viewer, Version 3 (GPL-3.0)] was used to calculate CT density measurements of the left lungs. Regional CT Hounsfield units (HU) were acquired by Fiji (Image) 1.52p, National Institutes of Health, USA, [http://image.nih.gov/ij](http://image.nih.gov/ij) by manually creating a mask for every lung to remove background noise and sequentially using the analysis ‘Histogram’ to reveal the distribution of HU over the grafts. HU were considered as follows: portions between $-1000$ and $-900$ HU as ‘over-inflated’, between $-900$ to $-500$ as ‘well-aerated’, $-500$ and $-100$ HU ‘poorly aerated’ and $-100$ and 0 HU ‘not aerated’ [15].

**Statistics**

Data were visualized as median [25–75% interquartile range (IQR)] unless otherwise stated. Since all data were not normality distributed, non-parametric tests were applied. Mann–Whitney test was used to compare both groups and Friedman test for repeated measurements (Sidak test for post hoc analysis). All analyses were performed with Graphpad® Prism 7.04 (GraphPad Software Inc, La Jolla, CA, USA). $p$ values less than 0.05 were considered significant.

**Results**

**Baseline characteristics**

Baseline characteristics are listed in Table 1 and did not differ between both groups. Study design is illustrated in Fig. 2.

**EVLP physiology**

Inspiratory flow of VCV was 60 L/min, median inspiratory flow of FCV was 3 (3–4) L/min. Perfusate flows were not different between both groups VCV 1.51 (1.51–1.52) L/min vs. FCV 1.50 (1.49–1.51) L/min ($p = 0.77$). The median delivered tidal volumes were 269 (252–278) mL in VCV, and 271 (249–286) mL in FCV ($p > 0.99$). Median PAP and median LAP were similar between both groups (VCV: 14.3 (13.6–14.7) mmHg vs. FCV: 15.6 (14.7–16.0) mmHg; $p = 0.21$; and VCV: 3.2 (2.9–3.5) mmHg vs. FCV: 3.3 (2.9–3.5) mmHg; $p = 0.87$), respectively.

Results of the 6-h EVLP interval are visualized in Table 2 and Fig. 3. PVR was comparable between VCV and FCV ($p = 0.52$). Oxygenation ratio (PaO$_2$/FiO$_2$ (P/F) ratio) was significantly higher in FCV compared to VCV ($p = 0.01$). Sidak post hoc analysis revealed significant differences at time points 1, 3 and 4 h during EVLP ($p = 0.04$, $p < 0.01$ and $p < 0.05$, respectively). Because of the difference in measurement of compliance, no conclusions can be withdrawn comparing the absolute values between both groups. In addition to the absolute values, we analyzed the difference ($\Delta$) of each physiological parameter between 6 and 1 h of EVLP. No differences were detected in $\Delta$PVR ($p = 0.53$) and $\Delta$ oxygenation ($p = 0.32$). The $\Delta$ compliance between 6 and 1 h of EVLP was significantly greater in VCV compared to FCV ($p = 0.03$).

Weight gain between pre- and post-EVLP was comparable between both groups [VCV 195 g (−27 to 230) and FCV 37 g (−53 to 130); $p = 0.38$].
Analyses of the BAL fluid did not reveal any difference in median total cell count ($\times 10^9$ cells/mL) (2.9 (2.7–3.0) in VCV vs. 3.0 (2.5–3.4) in FCV; $p = 0.97$) or in differential cell...
count: number of macrophages (90% (88–94) in VCV vs. 90% (87–91) in FCV; \( p = 0.62 \)) and neutrophils [10% (6–12) in VCV vs. 10% (9–13) in FCV; \( p = 0.62 \)] were similar. Cell viability was comparable between both groups [38% (25–40) in VCV vs. 46% (36–66) in FCV; \( p = 0.26 \)].

### W/D ratio, histology and CT analyses

Results of W/D ratio, CT imaging and pathology are visualized in Fig. 4. Macroscopic aspects of the grafts after EVLP are illustrated in Fig. 4d. Representative CT images of both groups are visualized in Fig. 4e and pathology images are shown in Fig. 4f. In line herewith is the absence of difference in W/D ratios between groups (\( p = 0.16 \)). Total CT density measurements tended to be lower in FCV (\( p = 0.05 \); Fig. 4e). There was no difference in composite pathology score (\( p = 0.16 \); Table 3). Further analyses of HU distribution throughout the lungs are illustrated in Fig. 5. No differences were observed in

---

**Table 2 Peak, plateau pressure and perfusate lactate, pCO₂ and pH values between both groups**

| Metric                       | VCV (n = 7) median with IQR | FCV (n = 7) median with IQR | \( p \) value (Mann–Whitney) |
|------------------------------|-----------------------------|-----------------------------|-----------------------------|
| Peak pressures at 6 h EVLP   | 20 (17–25) cm H₂O          | 22 (20–27) cm H₂O          | \( p = 0.51 \)               |
| Plateau pressures at 6 h EVLP| 8 (8–9) cm H₂O             | –                          | \( p = 0.62 \)               |
| Intratracheal pressures at 6 h EVLP | –            | 22 (20–27) cm H₂O         | –                          |
| pCO₂ at 6 h EVLP ‘inflow’ (pulmonary artery) | 42 (40–43) mmHg | 41 (39–44) mmHg          | \( p > 0.99 \)              |
| pCO₂ at 6 h EVLP ‘outflow’ (left atrium) | 36 (35–39) mmHg | 37 (36–39) mmHg          | \( p = 0.38 \)              |
| pH at 6 h EVLP outflow       | 7.40 (7.39–7.41)           | 7.38 (7.37–7.40)           | \( p = 0.23 \)              |
| Lactate at 6 h EVLP outflow  | 12 (11–13) mg/dL           | 13 (13–14) mg/dL           | \( p = 0.18 \)              |

VCV volume-controlled ventilation, FCV flow-controlled ventilation, IQR interquartile range
the proportion of $-1000$ and $-900$ HU (‘over-inflated’) when comparing VCV and FCV ($p > 0.99$), nor between $-500$ and $-100$ HU (‘poorly aerated’ $p = 0.94$). A higher proportion of HU between $-900$ and $-500$ was observed in FCV compared to VCV (‘well-aerated’, $p = 0.01$). The proportion between $-100$ and 0 HU tended to be lower in FCV compared to VCV (‘non-aerated’ $p = 0.06$).

Discussion

This is the first study evaluating the effects of the novel ventilation method FCV in a large animal EVLP model. We demonstrated that FCV as ventilation strategy during EVLP is feasible and might improve graft function. When comparing FCV and VCV, oxygenation was superior in FCV and with a better maintenance of recalculated compliance over
6 h in FCV. In addition, the proportion ‘well-aerated’ lung parts (HU between -900 and -500) was higher in FCV. These parameters suggest an improved preservation of alveolar recruitment in FCV compared to conventional VCV. Further studies with longer EVLP times and transplantation models are necessary to confirm that FCV better preserves lung grafts and limits injury during EVLP preservation.

However, in this study, there was no difference in total lung injury. Extravascular water content, measured by W/D ratios, weight gain after EVLP and cellular differentiation of BAL samples did not differ between FCV and VCV. Furthermore, pathology scores were similar between both groups. These findings suggest that in these 6 h of EVLP, FCV did not affect extravascular water accumulation nor inflammatory responses.

The main difference between FCV and VCV is that in FCV, both inspiration and expiration phase are controlled, whereas in VCV, expiration occurs by elastic recoil of the lungs [16]. The constant flow during expiration in FCV, might prevent sudden alveolar collapse and might delay complete emptying of the alveoli resulting in improved recruitment (less atelectasis) and better oxygenation [13]. The reduced number of collapsed alveoli decreased the risk for local overdistention alveoli (less volutrauma), and thereby potentially decreases an important driver of VILI.

Since atelectrauma and volutrauma both trigger inflammatory responses and contribute synergistically to VILI [10], development of new ventilation strategies which avoid these traumatic events are crucial—especially in donor lungs which are already at risk of developing ischemia–reperfusion lesions because of donor-related injury, emerging during ventilation after brain insult and further during procurement and preservation on ice.

The potential lung protective effects of controlling the expiratory flow were investigated by Goebel et al. [16]. Expiratory flow was slowed down by applying a flow restrainer in the expiratory limb of the circuit of a conventional ventilator and this flow-controlled expiration (so called “FLEX”) was evaluated in porcine ARDS lungs. Improved lung compliance and reduced lung pathology in the “FLEX” group compared
to the conventionally ventilated group could be demonstrated. In addition, in healthy
and injured pig lungs, “FLEX” was shown to lead to a more homogenous ventilation and
shifting ventilation from ventral to dorsal parts [17]. Similar results were observed in
healthy subjects [18].

FCV is a more sophisticated way to control the expiratory flow, as (in contrast to
“FLEX”) it is linearized by actively regulating the egress of gas by controlled suctioning
instead of using a passive flow restrainer. This allows to achieve a preferable I:E ratio of
1:1 without the risk of undesired auto-PEEP to build up [13]. Furthermore, the FCV ven-
tilator has been designed and built to also meet the requirements of a measurement tool
allowing to study dynamic lung mechanics by the combination of minimized technical
dead space, precise volumetry (by mass flowmetry) and continuous intratracheal pres-
dure measurement. However, in our study, these features were not applied to individual-
ize the settings.

To our knowledge, this is the first study that uses the innovative FCV mode in an
ex vivo setting. Other groups investigated the effects of FCV in vivo. Two previous stud-
ies [13, 19] compared the effects of FCV versus VCV in healthy pigs or ARDS pigs. Both
studies demonstrated a significantly higher PaO2 during FCV at a lower minute volume.
Additionally, CT scans of FCV lungs revealed an increased proportion of normally
aerated lung volume and a reduced area of poorly or non-aerated lung parts compared
to VCV [13, 19], which is in line with the observations in our study. More recently, a ran-
donized controlled trial was published, comparing FCV with VCV in patients under-
going laryngeal surgery [20]. FCV resulted in improved lung aeration and increased
respiratory system compliance, while using a lower inspiratory plateau pressure com-
pared to VCV.

Next to volutrauma, barotrauma and atelectrauma, mechanical power, has become
acknowledged as a causal factor for development of VILI [21]. Excess of mechanical
power applied to lungs has been shown to have deleterious effects [21, 22].

As explained, FCV is based on generating a constant, linearized low flow into and out
of the lungs, resulting in slow, continuous, linear increases and decreases of intratra-
cheal and thus intrapulmonary pressures that are just enough to establish mechanical
ventilation with efficient gas exchange. The sudden alveolar pressure drop during uncon-
trolled passive expiration in conventional ventilation is prevented. In other words, the
amount of energy generated by the FCV ventilator is just enough to establish sufficient
gas exchange. Thereby, the impact on the lung tissue by dissipated energy is kept to a
minimum, enabling ventilation with a markedly reduced risk of lung damage. Recently,
clear theoretical evidence was provided for lower energy dissipation in the lungs by
FCV as compared to VCV or PCV [23]. This theory was further validated in a patient,
showing that the energy dissipation was just 0.17 J/L, which is even lower than values
reported for spontaneous breathing (0.2–0.7 J/L) [24]. In the current study, dissipated
energy during VCV and FCV was not calculated, but the described lower energy dissipa-
tion by FCV may have contributed to the observed positive effects.

Limitations of this study might have been the large animal model with 2 h of warm
ischemic time, followed by 6 h of EVLP. One could argue that this is limited in time, but
physiologic differences already became visible during this ventilation strategy after 1 h
(increased oxygenation in FCV). This increase in oxygenation might be contributed by
increased $P_{\text{trach}}$ in FCV. Compliance was recalculated differently between both groups and therefore complicates true comparison. Therefore, we reported the decline of compliance in time within each group as an important parameter of onset of lung injury. We did not record and compare mean AWP between both groups. In this study, we applied standard settings of each ventilation mode. We cannot exclude a higher mean AWP in FCV (also because of differences in I:E ratio) which might have contributed to our findings. Further research should clarify the influence of mean AWP between both ventilation modes before generalization of these results. In addition, regional CT-graphic changes were already visible after 6 h of EVLP. These scans were taken at the end of every experiment. Whether a prolonged period of EVLP may have revealed differences in total lung injury remains to be studied. Though, the improved alveolar recruitment and maintenance of lung mechanics (compliance) indicate that FCV could lead to less VILI and enable longer perfusion times.

**Conclusions**

In conclusion, we observed that FCV leads to increased aerated regions in the lung during EVLP compared to conventional VCV in a large animal model. Future studies are necessary to confirm that FCV better preserves lung grafts and limits injury during EVLP preservation. This opens the perspective of a possible extension of EVLP duration and better preservation of graft quality during EVLP.

**Abbreviations**

BAL: Broncho-alveolar lavage; CO₂: Carbon dioxide; CT: Computed tomography; EVLP: Ex vivo lung perfusion; FCV: Flow-controlled ventilation; FiO₂: Fraction of inspired oxygen; HU: Hounsfield unit; I:E: Inspiratory time-to-expiratory ratio; LAP: Left atrial pressure; LTx: Lung transplantation; P/F: $\text{PaO}_2$/FiO₂; PAP: Pulmonary artery pressure; PCV: Pressure-controlled ventilation; PEEP: Positive end-expiratory pressure; PVR: Pulmonary vascular resistance; RR: Respiratory rate; TV: Tidal volume; VCV: Volume-controlled ventilation; VILI: Ventilator-induced lung injury; W/D ratio: Wet-to-dry weight ratio.

**Acknowledgements**

The authors would like to sincerely thank Mrs. Nicole Jannis, Miss. Caroline Lauwers, Mr. Tom Lauwers, Miss. Jozefien Van Ranst, Mr. Cedric Luyten and Miss. Mélanie Nolmans for their technical assistance during the animal experiments. Miss Natalie Alderson is thanked for her assistance in pathology processing. Mr. Walter Coudyzer is greatly thanked for his technical support regarding the CT imaging. Prof. Dr. Marc Van de Velde is greatly thanked for his support throughout the whole study. Dr. José van der Hoorn, Dr. Lena Bottinger and Prof. Dr. Dietmar Enk are greatly thanking for their critical review of this manuscript.

**Authors’ contributions**

SO, MO, TH, JK, BS, AS, and AVH performed the research work and data collection. SO interpreted the BAL samples and performed statistical analysis. AVS scored pathology biopsies. SO, APN, BMV, DEV, GMV, RV, SEV and JAV contributed to the concept and design of the study. All authors read and approved the final article.

**Funding**

This study received financial support from the KULeuven, Leuven, Belgium (C24/18/073), and the Research Grant Program of the Society for Anesthesia and Resuscitation of Belgium (SARB). R.V. is a clinical research fellow of the Research Foundation FWO. The FCV ventilator (Evone®) and required disposables were provided by Ventinova Medical, Eindhoven, The Netherlands. The OCS™ solution was kindly offered by Transmedics (Andover, MA, USA) and the albumin solution used for the EVLP perfusate was donated by CAF-DCF (Brussels, Belgium). This did not affect the study.

**Availability of data and materials**

The dataset used and analyzed during the current study is available from the corresponding author on reasonable request.

**Ethics approval and consent to participate**

This large animal study was conducted after approval by the Ethics Committee of the KULeuven (P145/2016). Each animal received humane care conforming to the “Principles of Laboratory Animal Care”, formulated by the National Society for Medical Research and “Guide for the Care and Use of Laboratory Animals”, prepared by the Institute of Laboratory Animal Resources and reported by the National Institutes of Health, USA (NIH Publication No. 86-23, revised 1996).

**Consent for publication**

Not applicable.
Competing interests
The authors declare that they have no competing interests.

Author details
1 Unit of Anesthesiology and Algology, Department of Cardiovascular Sciences, Katholieke Universiteit Leuven, Leuven, Belgium. 2 Department of Anesthesiology, University Hospitals Leuven, Herestraat 49, 3000 Leuven, Belgium. 3 Leuven Lung Transplant Group, Katholieke Universiteit Leuven, Leuven, Belgium. 4 Department of Thoracic Surgery, University Hospitals Leuven, Leuven, Belgium. 5 Laboratory of Respiratory Diseases and Thoracic Surgery (BREATHE), Department of Chronic Diseases, Metabolism and Ageing (CHROMMETA), Katholieke Universiteit Leuven, Leuven, Belgium. 6 Department of Pneumology, University Hospitals Leuven, Leuven, Belgium. 7 Department of Pathology, University Hospitals Leuven, Leuven, Belgium. 8 Department of Radiology, University Hospitals Leuven, Leuven, Belgium.

Received: 18 June 2020 Accepted: 17 November 2020

Published online: 25 November 2020

References
1. Venuta F, Van Raemdonck D (2017) History of lung transplantation. J Thorac Dis 9(12):S458–S471
2. Van Raemdonck D, Neyrinck A, Cypel M, Keshavjee S (2015) Ex-vivo lung perfusion. Transpl Int 28(6):643–656
3. Warnecke G, Van Raemdonck D, Smith MA, Massard G, Kukreja J, Rea F et al (2018) Normothermic ex-vivo preservation with the portable Organ Care System Lung device, for bilateral lung transplantation (INSPIRE): a randomised, open-label, non-inferiority, phase 3 study. Lancet Respir Med 6(5):357–367
4. Van Raemdonck D, Rega F, Rex S, Neyrinck A (2018) Machine perfusion of thoracic organs. J Thorac Dis 10(S8):S910–S923
5. Yeung JC, Zamel R, Klement W, Bai X-H, Machuca TN, Waddell TK et al (2018) Towards donor lung recovery-gene expression changes during ex vivo lung perfusion of human lungs. Am J Transplant 18(6):1518–1526
6. Becker S, Steinmeyer J, Avsar M, Hoffker F, Salman J, Haverich A et al (2016) Evaluating acellular versus cellular perfusate composition during prolonged ex vivo lung perfusion after initial cold ischaemia for 24 hours. Transplant Int 29(1):88–97
7. Iskender I, Cypel T, Arns S, Trinkwitz M, Fehlings S, Yamada Y et al (2018) Cytokine filtration modulates pulmonary metabolism and edema formation during ex vivo lung perfusion. J Heart Lung Transplant 37(2):283–291
8. Bucelho MT, Himmatt S, Stewart CJ, Hatami S, Dromparis P, Adam BA et al (2019) Continuous hemodialysis does not improve graft function during ex vivo lung perfusion over 24 hours. Transplant Proc 51(6):2022–2028
9. Ordies S, Frick AE, Claes S, Schols D, Verleden SE, Van Raemdonck DE et al (2019) Prone positioning during ex vivo lung perfusion influences regional edema accumulation. J Surg Res 239(16):300–308
10. Bates JHT, Smith BJ (2018) Ventilator-induced lung injury and lung mechanics. Annu Rev Med 69:378–378
11. Betler J, Malhotra A, Thompson B (2016) Ventilator-induced lung injury. Clin Chest Med 37(4):633–646
12. Slutsy AS, Raniere VM (2013) Ventilator-induced lung injury. N Engl J Med 369(22):2126–2136
13. Schmidt J, Wenzel C, Mahn M, Spassov S, Schmitz HC, Borgmann S et al (2018) Improved lung recruitment and oxygenation during mandatory ventilation with a new expiratory ventilation assistance device. Eur J Anaesthesiol 35(10):736–744
14. Martens A, Ordies S, Vanaudenaerde BM, Verleden SE, Van Raemdonck DE et al (2017) Immunoregulatory effects of multipotent adult progenitor cells in a porcine ex vivo lung perfusion model. Stem Cell Res Ther 8(11):159
15. Gattinoni L, Caron P, Pelosi P, Goodman LR (2001) What has computed tomography taught us about the acute respiratory distress syndrome? Am J Respir Crit Care Med 164(9):1701–1711
16. Goebel U, Haberstroh J, Foerster K, Dassow C, Priebe HJ, Guttmann J et al (2014) Flow-controlled expiration: a novel ventilation mode to attenuate experimental porcine lung injury. Br J Anaesth 113(3):474–483
17. Borgmann S, Schmidt J, Goebel U, Haberstroh J, Guttmann J, Schumann S (2018) Dorsal recruitment with flow-controlled expiration (FLEX): a crossover study in mechanically ventilated lung-healthy and lung-injured pigs. Crit Care 22(1):1–9
18. Wirth S, Springer S, Spaeth J, Borgmann S, Goebel U, Schumann S (2017) Application of the novel ventilation mode Flow-Controlled Expiration (FLEX): a crossover proof-of-principle study in lung-healthy patients. Anesth Analg 125(4):1246–1252
19. Schmidt J, Wenzel C, Spassov S, Borgmann S, Lin Z, Wollborn J et al (2020) Flow-controlled ventilation attenuates lung injury in a porcine model of acute respiratory distress syndrome: a preclinical randomized controlled study. Crit Care. 48(3):e241
20. Schmidt J, Günter F, Weber J, Kehm V, Pfeiffer J, Becker C et al (2019) Glottic visibility for laryngeal surgery: tritube vs. microlaryngeal tube: a randomised controlled trial. Eur J Anaesthesiol. 36(12):963–971
21. Tonetti T, Vasques F, Rapetti F, Maiolo G, Collino F, Romiti F et al (2017) A driving pressure and mechanical power: new targets for VILI prevention. Ann Transl Med 5(14):1–10
22. Cressoni M, Gotti M, Chiurazzi C, Massari D, Alemi I, Amini M et al (2016) Mechanical power and development of ventilator-induced lung injury. Anesthesiology 124(5):1100–1108
23. Barnes T, van Asseldonk D, Enk D (2018) Minimisation of dissipated energy in the airways during mechanical ventilation by using constant inspiratory and expiratory flows—flow-controlled ventilation (FCV). Med Hypotheses. 121(July):167–176. https://doi.org/10.1016/j.mehy.2018.09.038
24. Barnes T, Enk D (2019) Ventilation for low dissipated energy achieved using flow control during both inspiration and expiration. Trends Anaesth Crit Care 24:5–12

Publisher's Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.