Pulmonary Effects of Adjusting Tidal Volume to Actual or Ideal Body Weight in Ventilated Obese Mice

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Obese patients could be more susceptible to mechanical ventilation (MV)-induced lung injury than non-obese patients due to weight-dependent changes in lung properties. The aim of this study was therefore to evaluate the pulmonary effects of 2 hours low VT MV in a diet-induced obese mice model, with VT calculated on either the actual body weight (VTaw) or the ideal body weight (VTiw). First, we hypothesized that a MV with VTaw would be associated with altered lung mechanics and an increased lung inflammation. Second, we hypothesized that a MV with a VTiw would preserve lung mechanics and limit lung inflammation. We analyzed lung mechanics and inflammation using bronchoalveolar lavage (BAL) cell counts, flow cytometry tissue analysis and histology. Lung mechanics and inflammation were comparable in control and obese mice receiving VTiw. By contrast, obese mice receiving VTaw had significantly more alterations in lung mechanics, BAL cellularity and lung influx of monocytes as compared to control mice. Their monocyte expression of Gr1 and CD62L was also increased. Alveolar neutrophil infiltration was significantly increased in all obese mice as compared to controls. In conclusion, our findings suggest that protective MV with a VTaw is deleterious, with a marked alteration in lung mechanics and associated lung inflammation as compared to lean mice. With VTiw, lung mechanics and inflammation were close to that of control mice, except for an increased alveolar infiltrate of polymorphonuclear neutrophils. This inflammation might be attenuated by a blunted recruitment of inflammatory cells associated with obesity.

Over the last several decades, the worldwide prevalence of obesity has steadily risen, reaching 36% in the United States in 2010. Consequently, the number of obese inpatients requiring mechanical ventilation (MV), either in intensive care units or in the operating room, is increasing.

MV with excessive lung stress and strain may induce pulmonary adverse effects, called “ventilator-induced lung injury (VILI)”, due to barotrauma, volutrauma, atelectotrauma and biotrauma, leading to increased mortality in patients with the acute respiratory distress syndrome (ARDS). Some evidence has suggested that high tidal volume (VT) may also be deleterious for patients with a healthy lung, even for short periods of ventilation, as during general anesthesia. Recent guidelines recommend limiting VT to 6 to 8 mL/kg of ideal body weight...
Mass index (BMI) was also increased (0.40 [0.38–0.44] vs 0.31 [0.30–0.32] g/cm², measured at the beginning of the ventilation period showed that obese (OVT iw) and control (VC group) mice dis-

198 [188–226] in the two groups (196 [183–228] in NVO mice

g/mL in NVC mice,
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would preserve lung mechanics and limit lung inflammation as compared to a VT calculated on the actual body

Weight characteristics of the mice in the various groups. Mann-Whitney test and Kruskal-Wallis test were not significant between the control groups and between the obese groups, respectively. BMI: Body Mass Index.

|                      | NVC (n = 22) | VC (n = 24) | NVO (n = 21) | OVTaw (n = 19) | OVTiw (n = 21) |
|----------------------|--------------|-------------|--------------|---------------|---------------|
| Weight (g)           | 31.0 [29.5–32.1] | 30.8 [29.2–32.2] | 40.8 [37.7–42.6] | 42.0 [39.3–48.3] | 41.2 [38.6–45.3] |
| BMI (g/cm²)          | 0.31 [0.30–0.33] | 0.31 [0.30–0.32] | 0.39 [0.37–0.41] | 0.43 [0.39–0.46] | 0.41 [0.38–0.44] |

Table 1. Weight characteristics of the mice in the various groups. Mann-Whitney test and Kruskal-Wallis test were not significant between the control groups and between the obese groups, respectively. BMI: Body Mass Index.

|                  | NVC (n = 9–11) | NVO (n = 9–11) | p   |
|------------------|----------------|---------------|-----|
| BAL cell count (cells/mL) | 80,000 [68,300–95,556] | 73,333 [64,444–88,889] | 0.60 |
| BAL neutrophil (cells/mL)  | 444 [0–683] | 0 [0–433] | 0.16 |
| Macrophages (10⁶ cells/lungs) | 519 [394–723] | 560 [499–1018] | 0.11 |
| Neutrophils (10⁶ cells/lungs) | 882 [453–2,083] | 1,374 [912–2,555] | 0.36 |
| Monocytes (10⁶ cells/lungs)  | 246 [115–636] | 375 [230–534] | 0.34 |
| Histologic inflammation score | 3.0 [2.5–7.5] | 9.0 [6.0–11.0] | 0.08 |

Table 2. Characteristics of pulmonary inflammation in control and obese mice before ventilation. Pulmonary inflammation was characterized by BAL cell counts, number of inflammatory cells counted by flow cytometry and a histologic inflammation score. BAL: Bronchoalveolar lavage.

Results

Diet-induced obesity model. After the 12-week feeding period, the weight and BMI of all mice fed a high-fat diet (non-ventilated obese (NVO) + obese mice ventilated with VT based on actual weight (OVTaw) + obese mice ventilated with VT based on ideal weight (OVTiw)) was higher than that of mice fed a low-fat diet (non-ventilated controls (NVC) + ventilated controls (VC)): 41.1 [38.6–45.9] g vs 30.9 [29.5–32.1] g, p < 0.0001. Their body mass index (BMI) was also increased (0.40 [0.38–0.44] vs 0.31 [0.30–0.32] g/cm², p < 0.0001) (Table 1).

Moreover, obese mice might be more susceptible to VILI, even when ventilated with a low VT, because of both mechanical and biological characteristics of this condition. Firstly, obese individuals have a lower thoraco-pulmonary compliance because of fatty infiltration of the chest, which induces increased airway pressures, and their lungs may be not uniformly inflated during MV because of greater basal atelectasis. Secondly, obesity is associated with a baseline low-grade inflammatory status, which might increase vulnerability to bio-

trauma. Thus, low serum adiponectin levels have been shown to promote acute lung injury in obese mice. It can therefore be hypothesized that obese subjects may be more prone to MV-associated lung injuries than non-obese subjects, with low VT ventilation calculated on actual weight or even calculated on IBW. However, to the best of our knowledge, this hypothesis has not been previously investigated despite the important implications in terms of public health given the continuously growing prevalence of obesity.

The aim of this study was therefore to evaluate the pulmonary effects of low VT MV in a diet-induced obese mouse model, with VT calculated on either the actual body weight or the ideal body weight. First, we hypothesized that a MV with a VT calculated on the actual body weight would be associated with altered lung mechanics and an increased lung inflammation. Second, we hypothesized that a MV with a VT calculated on the ideal body weight would preserve lung mechanics and limit lung inflammation as compared to a VT calculated on the actual body weight. Some of our results have been previously reported in abstract form.

Baseline characteristics of obese and control mice. Pulmonary inflammation, assessed by BAL cell counts, histologic score and inflammatory cells in lung tissue quantified by flow cytometry, did not differ between NVO and NVC mice (Table 2). Alveolar permeability, assessed by the BAL protein concentration, was also similar in the two groups (196 [183–228] in NVO mice vs 198 [188–226] µg/mL in NVC mice, p = 0.84). Lung mechanics measured at the beginning of the ventilation period showed that obese (OVTiw) and control (VC group) mice displayed similar compliance (125 [115–131] vs 123 [111–127] ml/cmH₂O, p = 0.49). After recruitment maneuvers, obese mice (OVTiw and OVTaw) had higher peak and mean airway pressures than VC mice and OVTaw mice had higher peak and mean airway pressures than OVTiw mice (Fig. 1).

Lung mechanics during MV. Because the mean weight of VC mice was used to calculate the VT applied in OVTiw mice, the two groups had an identical VT of 0.241 [0.230–0.252] mL, which was significantly lower than the VT received by OVTaw mice (0.331 [0.304–0.376] mL, p < 0.0001). Lung mechanics slightly deteriorated

(IBW) in patients with ARDS, which tend to be extended to all critically ill patients and a VT < 10 mL/kg of IBW has also been recommended during anesthesia.

Despite the increasing number of ventilated obese patients, no specific study has been published concerning these characteristics of MV. Guidelines advocate the use of the IBW to calculate the VT, but the formulas are unfamiliar to medical doctors and are not fully integrated into current practice. Indeed, it has been recently demonstrated that obesity is a risk factor for being ventilated with a large VT (≥ 10 mL/kg of IBW), with an increased risk of secondary ARDS in obese ICU patients. Moreover, obese patients might be more susceptible to VILI, even when ventilated with a low VT, because of both mechanical and biological characteristics of this condition. Firstly, obese individuals have a lower thoraco-pulmonary compliance because of fatty infiltration of the chest, which induces increased airway pressures, and their lungs may be not uniformly inflated during MV because of greater basal atelectasis. Secondly, obesity is associated with a baseline low-grade inflammatory status, which might increase vulnerability to bio-trauma. Thus, low serum adiponectin levels have been shown to promote acute lung injury in obese mice. It can therefore be hypothesized that obese subjects may be more prone to MV-associated lung injuries than non-obese subjects, with low VT ventilation calculated on actual weight or even calculated on IBW. However, to the best of our knowledge, this hypothesis has not been previously investigated despite the important implications in terms of public health given the continuously growing prevalence of obesity.

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during the 2-hour MV period in the 3 ventilated groups, with an increase in airway pressures and a decrease in respiratory system compliance (Fig. 1). Deterioration of lung mechanics during the MV period was more pronounced in the OV Taw group, with a ΔH2-H0' higher for peak and mean airway pressures ($p = 0.00015$ vs VC and OV T iw) and lower for the respiratory system compliance ($p = 0.04$ vs VC and $p = 0.02$ vs OV T iw after corrections for multiple testing).

**BAL fluid cell counts and protein concentration.** Inflammatory cells in BAL fluid were almost exclusively macrophages (Fig. 2). The global cell count and the differential cell count of macrophages did not differ between VC and OV T iw groups. There was a higher global cell count in the OV Taw group as compared to the VC and NVO groups ($p = 0.0287$ and $0.0238$ after correction for multiple testing, respectively). These cells were macrophages and neutrophils, as macrophages and neutrophils cell counts were higher in the OV Taw group as compared to the NVO group ($p = 0.0455$ and $0.0434$, respectively) and tend to be higher as compared to the VC group ($p = 0.0959$ and $0.0602$ after correction for multiple testing, respectively). The BAL protein concentration did not differ between the three groups (Fig. 2).

**Inflammatory cell subpopulations and their activation by flow cytometry in lung tissue.** Both obese and control mice exposed to MV displayed higher monocyte and neutrophil counts in lung tissue, compared to non-ventilated mice (VC vs NVC, OV Taw vs NVO and OV T iw vs NVO comparisons). Similar macrophage counts were observed in the ventilated and non-ventilated groups. Within ventilated mice, we observed a higher monocyte count in lung tissue of OV Taw group (but not OV T iw animals) as compared to the VC group. Macrophage and neutrophil counts in lung tissue were not significantly different between the three ventilated groups (OV Taw, OV T iw and VC groups), but neutrophils also tended to be higher in OV Taw as compared to VC ($p = 0.0602$ after correction for multiple testing) (Fig. 3).

Concerning the degree of activation of the inflammatory cells, no difference in neutrophil CD11b and CD62L expression was observed between the 3 ventilated groups (Fig. 4d,e). In contrast, high monocyte activation was observed in OV Taw mice, as the expression of Gr-1 on monocytes was higher in the OV Taw group than in both the NVO and OV T iw groups. Moreover, the expression of CD62L on monocytes was higher in the OV Taw group than in the NVO group, whereas no difference was observed between the VC and NVC groups on one hand and between the OV T iw and NVO groups on the other hand (Fig. 4a,b). Interestingly, Gr-1 expression on monocytes was less intense in OV T iw mice than in VC mice (Fig. 4a).

**Quantification of lung tissue neutrophil infiltration.** Ventilated mice displayed higher neutrophil infiltration in the lungs compared to non-ventilated mice (VC vs NVC, OV Taw vs NVO and OV T iw vs NVO) (Fig. 5). This lung injury was particularly marked in obese mice, as the histologic score was higher in both the OV Taw and OV T iw groups compared to the VC group. No significant difference was observed between the OV Taw and OV T iw groups. Figure 5 represents the values of the histologic score and Fig. 6 shows representative sections of alveolar neutrophil infiltration.

**Discussion**

This study shows that ventilating obese mice with a $V_T$ calculated on their actual weight is deleterious, with a broader alteration of ventilatory mechanics, BAL total cell count, monocyte cell count and activation in lung tissue.

When $V_T$ was calculated using the actual weight, ventilatory mechanics were considerably impaired compared to ventilated controls, achieving the highest airway pressures and lowest compliance at H2. This cannot be explained *a priori* by more atelectasis, as these obese mice were subjected to the same level of PEEP. The
Figure 2. (a) Bronchoalveolar lavage (BAL) cell counts in the 5 groups of mice. (b,c) Number of macrophages (b) and neutrophils (c) in BAL. (d) BAL protein concentration. *p < 0.05 between the 2 groups after correction for multiple comparisons (n = 13–14 per group).

Figure 3. Numbers of neutrophils (a), monocytes (b) and macrophages (c) observed by flow cytometry in lung tissue in the 5 groups of mice. #p < 0.05; $p < 0.01 (n = 8–13 per group).
The most likely hypothesis is the presence of interstitial and alveolar edema, thereby inducing alveolar rigidity and decreased compliance, which may correspond to an early manifestation of ventilator-induced lung injury (VILI) in OVTaw mice. Although we did not demonstrate increased alveolar-capillary permeability in this group, it displayed increased BAL total cell count. Analysis on flow cytometry revealed an increase in recruited monocytes as compared to VC, and the number of neutrophils also tends to be higher. Moreover, monocytes of OVTaw displayed more intense L-selectin expression as compared to their non-ventilated counterparts, whereas no difference was observed between VC and OVT iw mice on one hand and VC and NVC on the other hand. Wilson et al.25 have previously shown that aggressive ventilation (35 mL/kg of VT in their model) and, to a lesser extent, protective ventilation, induced an influx of immature Gr-1 high monocytes from the bone marrow. These monocytes also overexpressed L-selectin, allowing them to migrate out of the bloodstream into the lung. Similar results were observed in our study, highlighting the harmful effect of a supposed “protective” VT calculated using the actual weight of obese mice. On histologic examination, we also demonstrated increased alveolar inflammation in OVTaw mice, with a higher neutrophil infiltration score compared to VC mice. The lack of significant increase in neutrophil by cytometry could be due to the short duration of the experiment (2 hours) that did not allow monocytes time to differentiate into neutrophils and/or the lack of infectious challenge.

Obese mice ventilated with a VT calculated using the ideal body weight had lung mechanics and inflammation parameters close to control mice except for a higher alveolar infiltration of neutrophils on histological score, which was similar to OVTaw, and a lower Gr-1 expression on monocyte. The decrease in most inflammatory parameters in the OVT iw group as compared to OVTaw may be ascribed to the reduction in VT with the use of ideal weight, but the higher neutrophil alveolar infiltration on histological score do not corroborate this hypothesis. An alternative hypothesis, supported by some publications, could be a role of obese phenotype in this decreased monocyte activation, as obese individuals appear to present a degree of immune deficiency, with delayed secretion of cytokines and recruitment of inflammatory cells in response to stimulation by a pathogen26,27, which could explain decreased activation of monocytes in OVT iw mice compared to VC mice, while alveolar infiltration by inflammatory cells was increased. Recent reports have highlighted abnormal diapedesis in obese mice exposed to inhalation of LPS28 or ovalbumin29 with increased influx of neutrophils and eosinophils in blood and lung tissue, but decreased BAL cell counts and cytokines. This phenomenon could also explain our

Figure 4. Gr-1 (a), CD62L (b) and CD11b (c) expression on pulmonary monocytes and CD62L (d) and CD11b (e) expression on pulmonary neutrophils expressed as mean fluorescent intensity (MFI). *p < 0.05, **p < 0.01 (n = 8–13 per group).
finding of excess neutrophils observed in the alveolar walls, but not in the BAL of OV_Taw mice. Overall, our findings may suggest that protective ventilation in obese subjects, even when VT is calculated using the IBW, induces an increased lung inflammation as compared to that observed in lean subjects, but this inflammation could be attenuated by a blunted recruitment of inflammatory cells associated with obesity.

This study presents several limitations. First, we used a short (2 hours) MV. However, a short period (2 to 3 hours) of low VT MV was shown to cause lung inflammation and edema in uninjured mice30, whereas a short period of high VT MV was shown to provide much more lung injury31–33. Second, we used a minimal PEEP (1.5 cmH2O). This might be criticized since numerous investigations encourage to use higher PEEP in the operating room, especially in obese patients34. However, other teams applied a similar low PEEP (below 2 cmH2O) during short MV in mice32,35,36. Here, with a PEEP settled at 1.5 cmH2O, we observed a slight deterioration of lung mechanics over the MV period. Indeed, the static compliance after 2 hours of MV was closed to that one recorded at time of connexion to the ventilator in both the VC and OV_Taw groups. In another work by our group, we showed that the dynamic compliance decreased below the H0 value after 3 hours of mechanical ventilation in uninjured mice37. Therefore, a 1.5 cmH2O PEEP looked suitable in such a low-VT short MV applied to uninjured mice. Third, our model of obesity was induced by diet, with a moderate weight gain, as compared to some genetic models38. This moderate weight gain may have precluded highlighting meaningful differences between obese and control groups. However, this model is much closer to clinical reality in humans and genetic models like ob/ob and db/db type could not be used in this study, as these animals present early weight gain that induces an abnormal development of the lungs, leading to a twofold lower lung volume than in control mice39. This discrepancy would have induced a major bias in the interpretation of the ventilatory mechanics. Fourth, we used the mean weight of control mice to calculate the ideal weight of obese mice. However, no formula is available to calculate IBW in mice. In addition, the naso-anal length of obese and control mice was similar in our study (data not shown). Fifth, some of our results (BAL cell count and flow cytometry neutrophils comparisons) were no longer statistically significant after correction for multiple testing, due to the low number of animals and the number of study groups. However, the corrected p values were “marginally significant”; overall, our results suggested that, with a short-term ventilation, the conjunction of obesity and a VT calculated on the actual weight induce
a significant alveolar inflammation. Finally, we did not use an infectious challenge like lipopolysaccharide, that could mimic the clinical scenario in intensive care where pneumonia is a major cause of mechanical ventilation. This lack of infectious challenge could in part explain the absence of enhanced inflammatory state in diet-induced obese mice at baseline before starting mechanical ventilation, like previously described in this model40. However, other studies have demonstrated the absence of overt lung inflammation in unchallenged obese mice41,42. Further experimental studies are needed to assess the exact impact of mechanical ventilation using the actual vs ideal body weight in terms of recruitment and migration of inflammatory cells (which could be blunted in obese individuals) and when using an infectious challenge like lipopolysaccharide.

Our study has some clinical implications. These data suggest systematically calculating the IBW of obese patients, both in intensive care and in the operative room, even for short periods of ventilation. However, further studies are required to assess the exact impact of protective mechanical ventilation on the IBW of obese subjects, as it appears to induce more marked pulmonary inflammation compared to controls, but the presence of secondary lesions must be determined. A deficiency in the recruitment and migration of inflammatory cells in obese subjects could also play a protective role against lung injury induced by inflammation secondary to mechanical ventilation. At present, clinical studies investigating the morbidity and mortality of ventilated obese subjects, often retrospective and comprising numerous biases, especially disparities between the management of obese and non-obese patients, have reported contradictory results43–46. Acute lung injury or ARDS may also have been over-diagnosed due to poor quality radiographs or undiagnosed atelectasis.

In conclusion, we herein demonstrate, for the first time, the harmful effect of a two hours ventilation strategy with tidal volume of 8 mL/kg calculated on the actual weight in obese mice. When the tidal volume was calculated using the ideal weight, lung mechanics and inflammation were close to that of control mice, except for an increased alveolar infiltrate of neutrophils on histology. Further studies are needed to determine the effect of protective ventilation based on IBW on the morbidity and mortality of these patients.

Material and Methods
Additional details on the methods are provided online (see supplementary information file, SIF).

Animal protocol. Five-week-old male C57/Bl6 mice were purchased from the Janvier laboratory (Le Genest-Saint-Isle, France) and were housed in our “small animal” facility. They received one week of adaptation with their usual food. At the age of 6 weeks, they were divided into two groups. One group (control group) received a diet in which 10% of the calories were derived from fat in the form of lard (D12450B, SSNIF, Soest, Germany), while the other group (obese group) was fed a diet in which 60% of the calories were derived from fat in the from of lard (D12492, SSNIF). Both groups were given food and water ad libitum and were exposed to a 12:12 h light-dark cycle. At the age of 18 weeks and after 12 weeks of the diet, animals were weighed and then fasted for 12 hours with free access to water before the experimentation. The body mass index (BMI) was calculated as previously described formula: BMI = weight (g)/(naso-anal length (cm))^2. Mice were then anesthetized with intraperitoneal pentobarbital (30 µg/g of body weight, Hospira, Meudon La Forêt, France) followed by continuous 1.5% isoflurane (Abbott, Rungis, France). The trachea was surgically exposed and the trachea was intubated orally under direct vision with a metal cannula (inner diameter of 1 mm, Harvard Apparatus, Les Ulis, France). The tracheal cannula was properly secured with surgical thread (Ethicon 3–0, Ethicon, Auneau, France) before connection to mechanical ventilator in order to avoid leaks.

Mechanical ventilation. After intubation, animals were left for 5 min in spontaneous breathing and half of the mice (obese and control) were sacrificed. The other half of were ventilated by means of a small-animal ventilator (flexiVent, Scireq, Montreal, Canada) as follows: V_T = 8 ml/kg of body weight, respiratory rate = 180/min, end-expiratory pressure = 1.5 cm H2O, and inspired fraction of oxygen = 0.4–0.6. This combination of V_T (8 ml/kg) and respiratory rate (180/min) has been shown to produce adequate minute ventilation and PaCO2 in mice45. Mechanical ventilation lasted 2 hours with continuous anesthesia maintained by 1.5% isoflurane and muscle paralysis using intraperitoneal pancuronium at the onset of the experiment (0.8 mg/kg of body weight, Organon, Puteaux, France) to ensure passive mechanical conditions. Body temperature was monitored using a rectal probe and was maintained at 36.5 °C with a blanket connected to a homeothermic regulator (Homoair-blanket system 50–7221 F, Harvard Apparatus, Les Ulis, France). Mice received one intraperitoneal warm fluid bolus (physiological serum) at the onset of the experiment (20 mL/kg of body weight). The dosage of all medications administered was calculated on actual weight. In ventilated obese mice, tidal volume was calculated either on actual weight or ideal weight. As no formula is available to calculate predicted body weight in mice, ideal weight was based on the mean weight of control mice (namely 31.2 g) (see SIF for details).

Experimental design. The experimental design included five groups: NVC (non-ventilated control mice), VC (ventilated control mice), NVO (non-ventilated obese mice), OV_Taw (ventilated obese mice with V_T calculated on the actual weight) and OV_Tiw (ventilated obese mice with V_T calculated on an ideal weight) (Fig. 7). The number of animals used in each experiment is detailed in the legend of all result table or figure.

Respiratory mechanics. Mean and peak airway pressures were recorded at initiation of mechanical ventilation and then hourly throughout the experimental period. Quasi-static compliance of the respiratory system was determined from a continuous pressure-volume curve performed at initiation of mechanical ventilation and at the end of MV.
red blood cells, the cell pellet was incubated for 5 min at room temperature with 5 mL of Gey’s solution (NH₄Cl and
inflated by tracheal injection of 4% formaldehyde at a pressure of 20 cmH₂O, fixed in 4% formaldehyde and finally
monary circulation was flushed by injection of 2 mL of saline into the right ventricle, and the lungs were then
imen collection procedures were used: For one half of the mice, immediately following thoracotomy, the pul -
(1500 rpm, 13 min at 4 °C), and cell-free supernatants were stored at

cell counts were performed on cytospin preparations (Cytospin 3; Shandon Scientific, Cheshire, UK) stained
counting grids (Hycor biomedical, Indianapolis, IN, USA). The cell pellet was diluted in saline, and differential
zymes for CD11b, CD11c, Gr-1 (Ly6C/G), F4/80 and CD62L (L-selectin) or appropriate isotype-matched controls.
ies for CD11b, CD11c, Gr-1 (Ly6C/G), F4/80 and CD62L (L-selectin) or appropriate isotype-matched controls.
ected as high CD11c and low CD11b cells. In order to differentiate and quantify mono -
cytometry analysis.

Specimen collection. After 2 hours of MV, mice were sacrificed by exsanguination. Two different spec -
were cut and stained with hematoxylin, eosin and saffron (HES). An optical microscope (BX51 Olympus) with camera
Histologic examination. After formalin fixation and paraffin embedding, 5µm thick sections of the left lung
Bronchoalveolar lavage. The total cell count was determined for a fresh fluid specimen using slides with count -
Bronchoalveolar lavage fluid was centrifuged (1500 rpm, 13 min at 4 °C), and cell-free supernatants were stored at —80 °C for subsequent assessment of protein content (using a colorimetric BCA assay, Thermo Scientific, Rockford, IL, USA).
Flow cytometric analysis. After performing BAL, a pulmonary circulation flush was performed as previously
described and both lungs were mechanically disrupted for flow cytometry analysis.

Figure 7. Design of experimental groups. (a) Mice were fed either a diet containing 10% fat (control mice) or
60% fat (obese mice) for 12 weeks. Non-ventilated control or obese mice (NVC and NVO) underwent general
anesthesia, were intubated and were sacrificed after 5 min of spontaneous ventilation. (b) Mice were fed the
same diet (10% or 60% fat). Ventilated control and obese mice underwent general anesthesia, were intubated
and were sacrificed after 2 hours of mechanical ventilation, with a tidal volume calculated on actual weight for
control mice (VC) or on either actual weight (OV-aw) or ideal weight (OV-iw) for obese mice.
Statistical analysis. Data were analyzed using Prism software version 5.0 (San Diego, CA, USA). Continuous data were expressed as median [IQ25–IQ75]. Statistical comparisons between two groups used the Mann-Whitney or the Wilcoxon test. Statistical comparisons between more than two groups used the Kruskal-Wallis test followed by a pairwise Mann-Whitney test with Benjamini-Hochberg correction for multiple testing. Two-tailed p values less than 0.05 were considered significant.

Ethical approval. All the experiments were performed in accordance with the official regulations of the French Ministry of Agriculture and the US National Institute of Health guidelines for the experimental use of animals, were approved by the animal institutional review board of the Paris-Est Creteil Val De Marne University, and were conducted in a specific “little animal” platform (Plateforme Exploration Fonctionelle du Petit Animal, INSERM U955, Creteil, France).

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E.G., A.R., M.D. and J.B. designed the study. E.G., A.R., S.K. and J.T.V.N. acquired the data. E.G., A.R., J.T.V.N., P.M., B.M., A.M.D., M.D. and J.B. analyzed and interpreted the data. All authors drafted the article, and approved the final version of the manuscript.

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