Stability of the animal model of oleic acid-induced acute lung injury*

Estabilidade do modelo animal de lesão pulmonar aguda induzida por ácido oleico

Eduardo Gaio, César Augusto de Melo e Silva, Flávio Brito, Marco Aurélio Pereira Firmino, Rodrigo Storck, Eduardo Freitas

Abstract

Objective: To evaluate the stability of hemodynamic, respiratory and gas exchange variables in an animal model of oleic acid-induced acute lung injury. Methods: This was an experimental study involving 10 mongrel dogs. The variables were measured at baseline, as well as at 30, 60, 90 and 120 min after the administration of oleic acid. In order to analyze repeated measurements, linear and quadratic effects were tested. Mixed linear models with diversified variance and covariance structures were used, depending on the variable studied. Results: We found that mean arterial blood pressure stabilized at 30 min, as did heart rate, pulmonary arterial pressure and pulmonary capillary pressure at 60 min. Respiratory rate, tidal volume, minute volume and respiratory work stabilized at 60 min. Regarding gas exchange variables, PaO$_2$, PaO$_2$/FiO$_2$ ratio and pulmonary shunt fraction stabilized at 30 min. The remaining variables maintained a continuous rise or fall. Conclusions: This oleic acid-induced acute lung injury model is stable for some of the variables tested, although stabilization occurs at different times. The respiratory and gas exchange variables stabilized at 30 min, whereas the hemodynamic variables stabilized at 60 min.

Keywords: Respiratory distress syndrome, adult; Models, animal; Statistical analysis.

Introduction

Acute lung injury (ALI) is a common entity, the evolution of which is occasionally rapid and usually severe. Its incidence varies by age bracket and has increased in recent years. Invariably, individuals with ALI require intensive care and mechanical ventilation. The study of the pathophysiology of ALI has advanced, and its therapeutic approach has gained prominence in the literature. Instabilities in biological variables, from the onset of ventilation to positive pressure, are well known, especially the decrease in cardiac output (Qt) due to offers of greater airway

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opening pressure, so that titration of mechanical ventilation parameters is necessary.\(^2\) Due to the complexity of its pathophysiology and to interactions in other systems, especially the cardiovascular system, ALI is the object of study in various fields of medical science. The animal models most commonly used in order to induce ALI are as follows: bronchoalveolar lavage with saline solution; intraperitoneal administration of paraquat; ischemia and reperfusion; and administration of oleic acid (OA) to the nervous system.\(^3\) Administration of OA to the nervous system was initially used as an animal model of pulmonary embolism.\(^4\) Subsequently, the OA-induced inflammatory injury was evaluated,\(^5\) this inflammatory injury being similar to that naturally found in lung injury.\(^6\) Venous infusion of OA causes capillary endothelial injury,\(^7\) has little effect on the relaxation/contraction properties of the endothelium,\(^8\) respects gravity-dependent areas\(^9\) and is dose-dependent.\(^10\) Gas exchange is known to worsen, although there is controversy regarding how long it takes the physiological variables to stabilize after OA administration.\(^11-14\)

Depending on the dose used, it is possible to keep the animal alive for up to seven days, although the evolution of the biological variables is debatable.\(^15\) In addition, the statistical methodology used in demonstrating the effects of time on the evolution of those variables is not the most appropriate.\(^16\) On the basis of this understanding, we studied, in function of time, the evolution of variables related to the hemodynamic and mechanical properties of the respiratory system, as well as that of gas exchange variables, in a model of OA-induced ALI. In addition, we tested the stability of those variables.

**Methods**

**Preparation of the animals**

This was an experimental study involving 10 male mongrel dogs, each weighing over 18 kg. The study was approved by the Animal Research Ethics Committee of the University of Brasília. The animals were initially sedated with i.v. thionembutal (30 mg/kg), were placed on the surgical table and were intubated with an 8.5-gauge orotracheal cannula. Two types of vascular access were installed in the right inguinal region: a venous access, for the administration of sedation and analgesia, and an arterial access, for the collection of blood for blood gas analysis and for the monitoring of mean arterial pressure (MAP). Venous access

<p>| Table 1 - Means of hemodynamic, respiratory and gas exchange variables during the experiment. |
|---------------------------------|----------------|----------------|----------------|----------------|----------------|</p>
<table>
<thead>
<tr>
<th><strong>Variable</strong></th>
<th><strong>Variable</strong></th>
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<th><strong>Variable</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>HR, bpm</td>
<td>142.4 ± 20.4</td>
<td>128.4 ± 15.4</td>
<td>151.6 ± 22.5</td>
<td>152.6 ± 22.4</td>
<td>157.0 ± 21.5</td>
</tr>
<tr>
<td>MAP, mmHg</td>
<td>125.5 ± 16.9</td>
<td>108.9 ± 14.2</td>
<td>96.9 ± 16.0</td>
<td>104.3 ± 7.3</td>
<td>102.0 ± 14.0</td>
</tr>
<tr>
<td>PAP, cmH(^2)/O</td>
<td>15.8 ± 5.5</td>
<td>15.8 ± 4.1</td>
<td>15.6 ± 5.1</td>
<td>13.9 ± 4.4</td>
<td>15.0 ± 5.5</td>
</tr>
<tr>
<td>PCP, cmH(^2)/O</td>
<td>5.2 ± 2.1</td>
<td>4.0 ± 1.9</td>
<td>3.3 ± 2.4</td>
<td>3.1 ± 1.1</td>
<td>3.1 ± 2.0</td>
</tr>
<tr>
<td>CI, L•min(^{-1})•m(^{-2})</td>
<td>8.25 ± 2.74</td>
<td>7.7 ± 2.1</td>
<td>7.2 ± 2.5</td>
<td>6.5 ± 2.8</td>
<td>6.2 ± 2.8</td>
</tr>
<tr>
<td>PVRI, dyn•s•cm(^{-6})•m(^{-2})</td>
<td>104.3 ± 27.3</td>
<td>123.2 ± 26.2</td>
<td>142.5 ± 53.7</td>
<td>145.2 ± 39.3</td>
<td>176.8 ± 80.4</td>
</tr>
<tr>
<td>RR, breaths/min</td>
<td>12 ± 6</td>
<td>100 ± 57</td>
<td>94 ± 51</td>
<td>86 ± 41</td>
<td>84 ± 39</td>
</tr>
<tr>
<td>(V_p), mL/kg</td>
<td>16.4 ± 4.1</td>
<td>6.6 ± 1.6</td>
<td>7.9 ± 1.6</td>
<td>8.1 ± 1.7</td>
<td>8.6 ± 2.2</td>
</tr>
<tr>
<td>(W, J)</td>
<td>2.351.5 ± 1.086.8</td>
<td>243.3 ± 162.0</td>
<td>401.0 ± 193.5</td>
<td>419.2 ± 119.2</td>
<td>425.3 ± 108.6</td>
</tr>
<tr>
<td>(Pa_{O_2}), mmHg</td>
<td>73.5 ± 13.5</td>
<td>45.2 ± 7.6</td>
<td>44.4 ± 11.6</td>
<td>43.5 ± 10.3</td>
<td>40.5 ± 9.7</td>
</tr>
<tr>
<td>(Pa_{CO_2}), mmHg</td>
<td>43.2 ± 7.4</td>
<td>49.6 ± 7.2</td>
<td>45.5 ± 12.3</td>
<td>39.8 ± 10.9</td>
<td>35.6 ± 8.5</td>
</tr>
<tr>
<td>(PA-a)(_{O_2}), mmHg</td>
<td>9.8 ± 7.3</td>
<td>28.1 ± 8.2</td>
<td>34.3 ± 14.9</td>
<td>42.4 ± 15.1</td>
<td>50.9 ± 13.6</td>
</tr>
<tr>
<td>(Pa_{O_2}/FiO_2), mmHg</td>
<td>349.8 ± 64.4</td>
<td>214.5 ± 36.3</td>
<td>210.8 ± 55.3</td>
<td>206.1 ± 48.2</td>
<td>190.5 ± 45.8</td>
</tr>
<tr>
<td>(Qs/Qt), %</td>
<td>34 ± 13</td>
<td>60 ± 10</td>
<td>59 ± 20</td>
<td>58 ± 10</td>
<td>60 ± 20</td>
</tr>
<tr>
<td>(DO_2), mL•min(^{-1})•m(^{-2})</td>
<td>1.237 ± 409.5</td>
<td>842.2 ± 207.4</td>
<td>740.3 ± 245.7</td>
<td>722.3 ± 157.2</td>
<td>652.4 ± 187.3</td>
</tr>
<tr>
<td>(VO_2), mL•min(^{-1})•m(^{-2})</td>
<td>213.6 ± 84.3</td>
<td>248.6 ± 53.6</td>
<td>223.3 ± 90.8</td>
<td>286.5 ± 119.7</td>
<td>244.4 ± 123.5</td>
</tr>
</tbody>
</table>

\(t_1\): baseline; \(t_2\): 30 min; \(t_3\): 60 min; \(t_4\): 90 min; \(t_5\): 120 min; HR: heart rate; MAP: mean arterial pressure; PAP: pulmonary artery pressure; PCP: pulmonary capillary pressure; CI: cardiac index; PVRI: pulmonary vascular resistance index; RR: respiratory rate; \(V_t\): tidal volume; W: respiratory work; \(PA-a\)\(_{O_2}\): alveolar-arterial oxygen gradient; \(Pa_{O_2}/FiO_2\): oxygen index; \(Qs/Qt\): pulmonary shunt fraction; \(DO_2\): oxygen delivery; and \(VO_2\): oxygen uptake. Results expressed as mean ± SD.
was maintained using 0.9% saline solution to prevent clots. Continuous infusions of midazolam (0.5 μg/kg/min) and fentanyl (5.0 μg/kg/min) were used for sedation and analgesia, those being the only continuous infusions used during the experiment. The right jugular was dissected so that a pulmonary artery catheter (model 131HF7; Edwards Lifesciences, Irvine, CA, USA) could be introduced. The location of the catheter was confirmed by the pulmonary artery pressure tracings seen on a multivariate monitor of biological signals (model DX 2010; Dixtal, Manaus, Brazil). Using this instrument, we collected mixed venous blood samples and measured Q_{t}, pulmonary artery pressure (PAP), pulmonary capillary pressure (PCP) and heart rate (HR). The variables calculated were the cardiac index (CI, calculated as CI = Q_{t}/m^{2}) and pulmonary vascular resistance index (PVRI, calculated as PVRI = (PAP - PCP) × 80/CI). The mechanical properties of the respiratory system and its components—lungs and rib cage—were evaluated using a Fleisch no. 0 pneumotachograph (Godart-Statham, Bilthoven, Holland) with a lateral outlet connected to the orotracheal cannula. We also used a differential pressure transducer (model PT5A; Grass Instruments, Quincy, MA, USA), which was connected to the outlets of the pneumotachograph in order to measure airway opening pressure. Another differential pressure transducer (same model) was connected to the esophageal cannula in order to measure esophageal pressure. The signals were filtered and amplified using a polygraph (model 7C; Grass Instruments) and subsequently sent to a biological signal conditioning module & 12-bit analog/digital converter (EMG System do Brasil, São José dos Campos, Brazil). The signals were registered and stored in a database created using the Windaq/Pro software (DATAQ Instruments, Akron, OH, USA), which was also used for the electronic integration of the flow signal in order to obtain the volume curve. The variables calculated were tidal volume (V_{f}), respiratory rate (RR), minute volume (MV, calculated as MV = V_{f} × RR), respiratory work (W = area of the pressure-volume curve) and respiratory work per minute (calculated as W/minute = W × RR). Arterial and mixed venous blood samples were collected for blood gas analysis and measurement of pulmonary shunt fraction, which is calculated on the basis of the systemic blood flow (Qs)/Qt ratio. The gas exchange variables measured were PaO_{2}, mixed venous oxygen pressure (PvO_{2}), PaCO_{2}, mixed venous carbon dioxide pressure (PvCO_{2}), mixed venous oxygen saturation (SvO_{2}) and pH. The variables calculated were as follows: bicarbonate; oxygen saturation (SaO_{2}); oxygen delivery (DO_{2}), which was calculated using the following formula: DO_{2} = (SaO_{2} × 1.36 × hemoglobin + PaO_{2} × 0.0039) × CI; oxygen uptake (VO_{2}), which was calculated using the following

Table 2 – Values of p for the linear and quadratic effects tested, as well as variance structures, for hemodynamic and respiratory variables.

<table>
<thead>
<tr>
<th>Type</th>
<th>Hemodynamic variables</th>
<th>Respiratory variables</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HR</td>
<td>MAP</td>
</tr>
<tr>
<td>Linear effect</td>
<td>0.0731</td>
<td>0.0001</td>
</tr>
<tr>
<td>Quadratic effect</td>
<td>0.8042</td>
<td>0.0067</td>
</tr>
<tr>
<td>t_{0} = t_{1} = t_{2} = t_{3}</td>
<td>0.0995</td>
<td>0.0004</td>
</tr>
<tr>
<td>t_{i} = t_{2} = t_{3} = t_{4}</td>
<td>0.0563</td>
<td>0.0718</td>
</tr>
<tr>
<td>t_{2} = t_{3} = t_{4}</td>
<td>0.9975</td>
<td>0.6100</td>
</tr>
<tr>
<td>Variances structure</td>
<td>CS</td>
<td>VCA</td>
</tr>
<tr>
<td>Stabilization/tendency</td>
<td>t_{2}</td>
<td>t_{1}</td>
</tr>
</tbody>
</table>

HR: heart rate; MAP: mean arterial pressure; PAP: pulmonary artery pressure; PCP: pulmonary capillary pressure; CI: cardiac index; PVRI: pulmonary vascular resistance index; RR: respiratory rate; V_{t}/kg: tidal volume per body mass; MV: minute volume; W: respiratory work; W/kg: respiratory work per minute; t_{0}: baseline; t_{1}: 30 min; t_{2}: 60 min; t_{3}: 90 min; t_{4}: 120 min; CS: compound symmetry; VCA: variance components autoregressive; HCS: heterogeneous compound symmetry; U: unstructured; and HA: heterogeneous autoregressive.
formula: \( \text{VO}_2 = ([\text{SaO}_2 - \text{SvO}_2] \times 1.36 \times \text{hemoglobin} + [\text{PaO}_2 - \text{PvO}_2] \times 0.0039) \times \text{IC} \); oxygen extraction ratio (\( \text{O}_2\text{ER} \)), which was calculated using the following formula: \( \text{O}_2\text{ER} = \text{VO}_2/\text{DO}_2 \); \( \text{PaO}_2/\text{FiO}_2 \) ratio; and \( \text{Qs/Qt} \).

**Experimental protocol**

After the values had stabilized, the variables described above were measured, and this was considered the baseline, designated time point 0 (\( t_0 \)). The dogs were maintained on spontaneous ventilation throughout the experiment, \( \text{FiO}_2 \) being 0.21. Five minutes after those variables were measured, ALI was induced by administration of OA (0.15 mg/kg for 10 min) through the proximal port of the pulmonary artery catheter. Hemodynamic, gas exchange and respiratory variables were assessed again at 30, 60, 90 and 120 min (\( t_1, t_2, t_3 \) and \( t_4 \), respectively) after the administration of OA. After the last variables had been assessed, the dogs were euthanized with a solution of KCl.

**Statistical methods**

The results were presented as mean ± SD. The null hypothesis was that the means of hemodynamic, respiratory and gas exchange variables were similar at \( t_0, t_1, t_2, t_3 \) and \( t_4 \). The possibility that the means of the variables at \( t_0 \) were different from those at the other time points and that the latter were similar to one another was the most plausible, which would signify stability after \( t_1 \) or \( t_2 \). The methodology used had a repeated measure design. In order to determine whether a polynomial, a linear or a quadratic function reasonably fit the data, an analysis of linear or quadratic tendency was performed. As a result, if the data fit only the linear effect, the mean of the variable would increase or decrease over time, and there would therefore be no stabilization. If solely the quadratic effect fit the data, the mean of the variable would increase or decrease up to a certain time point and would subsequently decrease or increase from that point onward. Therefore, there would be no stabilization. If the linear effect and the quadratic effect were present, the mean of the variable would markedly increase or decrease and, from a certain time point onward, could indicate stabilization. If none of the effects fit the data, the mean of the variable would remain constant over time. In parallel, the hypotheses of stabilization after \( t_1 \) and \( t_2 \) were tested. Mixed linear models with diversified variance and covariance structures were used, depending on the variable studied. The effect of time was evaluated using the generalized minimum squares method. For stabilization to occur at \( t_1 \), three conditions should be satisfied: the means at \( t_0, t_1, t_2, t_3 \) and \( t_4 \) must not be equal; the mean at \( t_0 \) must be different from the means at the other time points; and the means at \( t_1, t_2, t_3 \) and \( t_4 \) must not be different. For stabilization to occur at \( t_2 \), another three conditions should be met: the means at \( t_0, t_1, t_2, t_3 \) and \( t_4 \) must not be equal; the means at \( t_0 \) and \( t_1 \) must be different from the means at \( t_2, t_3 \) and \( t_4 \); and the means at \( t_2, t_3 \) and \( t_4 \) must not be different from one another. In order to reduce the alpha error, we chose to previously define the hypotheses and not to test all time points in relation to one another. The variables were evaluated for normal distribution using the Shapiro-Wilk test.
The level of statistical significance was set at 5% for all analyses. The SAS software (Statistical Analysis System, Cary, NC, USA) was used for all analyses described.

Results

Ten male mongrel dogs, with a mean weight of 26.7 ± 4.5 kg, were used. All of the animals survived until the end of the experiment. The values observed at the respective time points are presented in Table 1. For all variables, linear and quadratic effects were tested. For those variables that fit these effects, we tested the hypotheses that their means did not differ at $t_1$, $t_3$, $t_5$, and $t_7$, or at $t_2$, $t_4$, and $t_6$, which would signify stabilization at $t_1$ or at $t_2$, respectively. All variables, with the exception of DO$_2$, showed parametric distribution by the Shapiro-Wilk test.

Regarding hemodynamic variables, the analysis of linear and quadratic effects revealed that these effects were present for MAP, which stabilized at $t_1$. For HR, PAP and PCP, although no significance was found in the analysis of linear and quadratic effects, the values obtained at $t_2$, $t_4$, and $t_6$ ($p = 0.9975$, $p = 0.8669$ and $p = 0.8765$, respectively) were remarkably close, which suggests stabilization from $t_1$ onward. We observed that the CI and PVRI did not stabilize. The $p$ values for linear and quadratic effects and for the comparison of the time points for each of the hemodynamic variables studied, as well as the types of variance and covariance structure found and the time point at which stabilization or a tendency toward stabilization occurred, are shown in Table 2.

Regarding respiratory variables, a significant $p$ value was observed for RR, $V_{T}$ (mL/kg body weight), MV and W in the analysis of linear and quadratic effects. All of those variables stabilized at $t_1$. Only W/min increased indefinitely. The findings regarding these variables are shown in Table 2.

After the administration of OA, PaO$_2$, SaO$_2$, PaO$_2$/FiO$_2$ (Figure 1) and Qs/Qt (Figure 2) changed immediately, stabilizing at $t_1$. The alveolar-arterial oxygen gradient (P[A-a]O$_2$) stabilized at $t_2$. During the experiment, VO$_2$ did not change, whereas O$_2$ER and pH increased. In addition, PaCO$_2$ and bicarbonate decreased. None of the altered parameters stabilized. Since DO$_2$ did not present a normal distribution, it was transformed into a neperian logarithm. There was a continuous decline in DO$_2$ without stabilization. The effects tested, as well as the variance and covariance structures of these variables, are shown in Table 3.

Discussion

For decades, intravenous OA has been administered in order to mimic pulmonary embolism in animal models, and, since the 1970s, it has been administered in models of ALI. One group of authors conducted a study involving 22 dogs, who were kept alive for seven days in order to evaluate the various stages of the pathophysiology of ALI. That study revealed that
Qs/Qt and P(A-a)O₂ returned to values close to baseline values by the end of the experiment. There was significant worsening of these variables on day 1 and subsequent improvement until day 7. A more recent study[18] revealed the resolution of the edema on day 7 after infusion of OA. Another group of authors[19] showed the evolution of respiratory, hemodynamic and gas exchange variables but did not assess the stability of the model. Those authors studied 8 dogs for 2.5 h and compared the variables using ANOVA with repeated measures. There was an indefinite increase in PAP and PVRI. The increase in pulmonary compliance, PVRI and P(A-a)O₂ was designated as a recent event, as was the decrease in PaO₂, CI and MAP. The increase in PAP, Qs/Qt and PaCO₂ was treated as a late event. A study involving pigs[20] revealed stability of the model at 30 min for HR, PAP, Qs/Qt and dead space. In that study, ANOVA with repeated measures was also used in order to study the variables. In our study, MAP stabilized at t₁, whereas HR, PAP and PCP showed a tendency toward stabilization at t₂. Another group of authors[21] observed that those variables did not change during their study, which ensured the hemodynamic stability necessary for a model of ALI. The decrease and subsequent stabilization of the CI and gas exchange, as well as the increase in PVRI, are biologically plausible and accepted in the literature. It is likely that the decrease in myocardial contractility and the vascular lesion that are revealed by simple histological examination, despite there being no change in the wet-to-dry weight ratio, are responsible for the initial decrease in Qt.[22]

Another important factor for the decrease in the CI is the vasoplegia that occurs in this model. In our study, we observed a decrease in the CI using thermodilution and an increase in PVRI without stabilization—results that are discordant with those found in the literature. Other authors[23] demonstrated that, in rabbit lungs, it is not the total PVRI that changes but rather the segmental distribution of resistances. It is likely that our methodology was the reason for the discrepancies between our results and those of other authors, and that, if there is hemodynamic stabilization, it will occur after the stabilization of respiratory and gas exchange properties. In our study, it was clear that the increase in MV was due to the significant increase in RR, since Vₜ decreased. This determined the decrease in PaCO₂ and the nonstabilization of pH. We were unable to measure respiratory system compliance or resistance due to the high RR's observed. However, W decreased in each respiratory effort due to the significant decrease in Vₜ, although W/min increased throughout the experiment, suggesting a constant increase in compliance or airflow resistance and, therefore, in pulmonary impedance. When OA is administered, pulmonary compliance can increase up to three times in relation to its baseline value.[23]

It is also known that there is an increase in airway resistance, measured by using various methods, due to the accumulation of secretion.[23] One group of authors[24] measured the wet-to-dry weight ratio in pigs and observed that this ratio increased significantly in the group that received OA, compared with the control group, which suggests worsening of compliance. Using forced oscillation, the authors of another study[25] demonstrated an increase in airway opening pressure, although without stabilization for up to 140 min. The principal events responsible for the increase in compliance are the accumulation of liquid in the alveolar and interstitial space and the increase in surface tension. Since Qs/Qt seems to correlate with the severity of the lesion, the extravasation of intra-alveolar fluid tends to decrease during the experiment. With a 0.09 mL/kg dose of OA, shunt values are higher than 40%.[26] Using 0.08 mL/kg, other authors[27] registered shunt values of up to 35%. Those authors also demonstrated the decrease in and the stabilization of PaO₂ in the first 30 min, this condition persisting for 4 h. Another group of authors[27] found lower, basal fractions of Qs/Qt than those found in our study, approximately 15-20%. The dose used in the present study (0.15 mL/kg) explains the high Qs/Qt levels found. The 0.08 mL/kg dose of OA reduced the PaO₂/FiO₂ ratio from 451 to 139.[18]

Recent studies have relied on the potential stability of the OA-induced ALI model[28,29] All of the studies that assessed the stability of this animal model used ANOVA with repeated measures as the statistical methodology. There are two traditional methodologies for the study of longitudinal variables: univariate ANOVA with repeated measures and multivariate ANOVA with repeated measures. Unfortunately, these traditional methodologies are limited because,
a priori, they make assumptions that are false more often than not. Univariate analysis assumes that the variance and covariance structure of the events measured over time is symmetrical, which might not be true when the variances tend to increase over time and the correlations tend to decrease as the time intervals increase. The other alternative—multivariate analysis for repeated measures—only includes subjects with complete data, not allowing for potential losses during the study. In addition, these procedures estimate the tendencies of the group over time and pay little attention to individual changes. For these and other reasons, mixed models employing covariance have become popular as useful and powerful tools for the study of longitudinal data. In one interesting study, the researchers compared procedures for the analysis of repeated measures, highlighting the importance of the use of mixed linear models in a two-stage approach. In the first stage, the covariance structure is estimated. In the second stage, the estimate of covariance is replaced in the linear mixed model by the evaluation of the effect of time on the variable studied. In our study, we observed covariance structures that were symmetrical, heterogeneous, autoregressive and unstructured. The mixed linear model described a relationship among the values obtained for a certain variable at five time points as a function of their mean, of the inter-animal effect and of the intra-animal effect.

We are sure that, by increasing the number of animals, extending the study period and shortening the intervals between measurements of variables, we would gain a better understanding of the evolution and stability of hemodynamic, respiratory and gas exchange variables in OA-induced ALI. Forced oscillometry can assess respiratory mechanics in spontaneous ventilation of animals, extending the study period and shortening the intervals between measurements of a certain variable at five time points as a function of their mean, of the inter-animal effect and of the intra-animal effect.

In summary, we studied the stability of biological variables using a robust and modern statistical analysis and concluded that, in this model of OA-induced ALI, gas exchange and respiratory variables stabilize early, whereas hemodynamic variables stabilize later.

References

16. Grutjohan HP, van der Heijde RM, Wagenvoort CA, Wagenvoort N, Versprille A. Pulmonary vasoconstriction...


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