Mycoplasma infection and inflammatory effects on laboratory rats bred for experimental research

Infecção por micoplasma e efeitos inflamatórios em roedores criados em laboratório para pesquisa experimental

Infeción por micoplasma y efectos inflamatorios en roedores criados en laboratorio para investigación experimental

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ABSTRACT

Mycoplasma pulmonis is the agent of murine respiratory mycoplasmosis and is the most frequently isolated species from laboratory rats. One of the important consequences of its infection is the reduction of breeding rates and significant interferences in the experiments using these animals. Its experimental infection is also considered a model for studying human infection by M. pneumoniae. We have recently reported M. pulmonis infection rates of up to 75% in rodents bred in facilities for experimental research; however, its effect in the animals’ immune system was not evaluated. Therefore, the aim of this work was to investigate the effect of M. pulmonis in rats bred at the animal facility of University of Blumenau, by polymerase chain reaction and culture, in order to verify the viability of the use of these animals for experimentation. Inflammatory parameters, such as C-reactive protein (CRP) and alpha-1 acid glycoprotein (A1GP), as well as haematological parameters were determined from blood samples. It was observed a positivity rate of 80% of M. pulmonis infection. It was also noticed that mycoplasma infection was associated with increased CRP and A1GP levels, as well as blood monocytes. Considering that the possibility of mycoplasma infection is a well-known issue, and that the routine examinations and preventive measures have been well standardised internationally for many years, the findings of this study indicate that the use of the evaluated animals is unacceptable for experimental research. It is strongly advisable that laboratories which conduct this kind of research should be assured of the absence of mycoplasma infection in the animals were used.

Keyworks: Rats; Mycoplasmas; Lung Diseases; Hematology; Clinical Chemistry; Immunology.

INTRODUCTION

In the last few years, many advances have been made towards a better understanding of the pathogenicity of mollicutes (or mycoplasmas), especially regarding its interactions with the immune system, the demonstration of its ability to invade different cellular types, and the possibility that at least a few species may play a role in the development of AIDS1. These findings and the complete genome sequencing of several species have focused on increasing attention on mollicutes, the smallest microorganisms capable of self-replication.

The most pathogenic species in laboratory rodents is Mycoplasma pulmonis. The main disease caused by M. pulmonis is murine respiratory mycoplasmosis, the expression of which is influenced by a wide range of factors. It may cause significant morbidity and mortality, particularly in laboratory animals destined for long term experiments, as well as in animals suffering from a subclinical infection that may recrudesce by a laboratory manipulation2. Furthermore, the most important consequence of M. pulmonis infection, beyond its direct pathogenic effects to the animal, can be the consequence of its presence in the analysis of experimental data. For instance, it may affect ciliary cell
function, cellular kinetics\textsuperscript{3}, neurogenic inflammation\textsuperscript{1-5}, natural killer cell activity\textsuperscript{6,7}, local and systemic immune response\textsuperscript{2,8}, and induction of the production of several cytokines\textsuperscript{9}. Immune system activation by mollicutes induces the liberation of preinflammatory substances and polyclonal proliferation of B and T lymphocytes\textsuperscript{10}. Experimental \textit{M. pulmonis} infection is also considered the best model for studying human disease caused by \textit{M. pneumoniae}\textsuperscript{11}.

\textit{M. pulmonis} has been isolated from the urogenital tract of rats and mice\textsuperscript{2}. Another important consequence of this infection is a reduction of breeding rates, which may decrease by up to 50\%\textsuperscript{12}. \textit{M. pulmonis} might be also associated with arthritis in these animals, although a less supplicative disease than that caused by \textit{M. arthritidis}, the most frequent agent at that site\textsuperscript{2}. However, this disease may last for months, with immune complexes being detected in the joint tissues by immunofluorescence\textsuperscript{13,14}.

The efforts to reduce \textit{M. pulmonis} infection in laboratory rats have obtained a certain level of success, but have not been able to eliminate it. Prevalence studies from the 1980s still demonstrated rates of 5-20\%, even in well-controlled facilities\textsuperscript{2}. In spite of a lack of high scale studies in recent years, some groups have demonstrated that \textit{M. pulmonis} infection still persists. In the early 1990s, a significant mollicute infection rate was found in rodents of an animal facility in São Paulo State, Brazil, that did not have any microbiological barriers\textsuperscript{15}. More recently, in another animal facility in the same country, a \textit{M. pulmonis} infection rate of 65\% was found in laboratory rats, with colonisation of the personnel who was taking care of the facility\textsuperscript{16}. More recently, our group has described a \textit{M. pulmonis} infection rate of 75\% in Wistar rats bred at the University of Blumenau (FURB) facility\textsuperscript{7}.

In spite of the fact that mycoplasma infection is a well-known issue, and that the routine examinations and preventive measures have been well standardised internationally for many years, it is essential that laboratories be assured about the health of the animals used in experimental research. Unfortunately, the certification of animals and accreditation of facilities is not a reality in many countries. Therefore, the aim of this research was to evaluate the effect of \textit{M. pulmonis} infection in rats were bred at the animal facility of our institution regarding the levels of inflammatory proteins and peripheral blood cell types, as this information is unavailable in many research institutions that use this animal source.

\textbf{MATERIALS AND METHODS}

\textbf{ANIMALS AND SAMPLES}

In this study, 20 Wistar rats were bred at the animal facility of our institution were evaluated, equally distributed between males and females. This is a conventional animal facility, where the animals are bred in a clean area in polypropylene boxes with chrome-plated top cover, and feeder and drinker, but without microbiological barriers. Staff working with animals use equipments for individual safety when at work. Water and shavings are sterilised by autoclaving before entering into the clean area, and the shavings are disinfected with sodium hypochloride before being discarded. The animal facility supplies animals to meet teaching and research demand for students and professors of the health and biological areas. Periodical (annually) routine examinations for infectious diseases, as well as prophylaxis for parasites (semi-annually), are performed. The animal facility breeds rats and mice. For the experiments, animals were transferred to the local facility at the research laboratory or sampling. For mycoplasma detection, bronchoalveolar aspirates were obtained with 2 mL of culture media after euthanasia with thiopental (100 mg/kg), according to recommended standard procedures\textsuperscript{17}. For the evaluation of the inflammatory parameters, blood was obtained by cardiac puncture and 0.5 mL were dispensed in microtubes containing 50 \(\mu\)L of a ethylenediaminetetraacetic acid (EDTA) solution for white and red blood cell counts, and the remaining sample was dispensed in a dry microtube without anticoagulant for C-reactive protein (CRP) and alpha-1 acid glycoprotein (A1GP) determination. After coagulation of the samples without EDTA for 30 min at 37\(^\circ\) C, they were subjected to centrifugation at 2,500 rpm for 15 min, and then serum was separated to a new tube and maintained at -20\(^\circ\) C until analysis. The study was approved by the Committee of Ethics in Animal Research of the University of Blumenau under protocol no. 018/09 in August 26, 2009, and the animals were handled according to the Position Statements of American Association Laboratory Animal Service. It is worth noticing that from 2013 on it is recommended that in animals subjected to intraperitoneal anesthesia with barbiturates, when not possible intravenously, it must be mixed with lidocaine (10 mg/mL) or another local anesthetic\textsuperscript{19}.

\textbf{MYCOPLASMA CULTURE}

A 1 mL aliquot of the aspirates was immediately subjected to culture at 37\(^\circ\) C in a final volume of 2 mL of modified SP-4 liquid medium and plates with solid SP-4 medium (PPLO broth base [Difco], Tryptone [Oxoid], Peptone [Difco], L-arginine chloride, 1\% phenol red, VX supplemnt, 200 \(\mu\)g/mL penicillin, 20\% foetal calf serum, CRML 1066, 20\% yeast extract [Difco], and dextrose). Solid medium was prepared by adding 1\% Agar Noble (Difco). Samples were incubated for up to 30 days in aerobic conditions and observed daily for growth\textsuperscript{20}.

\textbf{DNA EXTRACTION AND PURIFICATION}

DNA from 1 mL aliquots of the bronchoalveolar samples was extracted by conventional lysis and phenol/ chloroform method. Samples in 1.5 mL microtubes were centrifuged for 15 min at 12,000 \(g\). The pellet was resuspended in 0.5 mL of lysis buffer (10 mM Tris, 1 mM EDTA, 0.5\% Triton X-100, pH 8.0) with 200 \(\mu\)g/mL Proteinase K, and incubated at 56\(^\circ\) C for 1 h, followed by 10 min of incubation at 100\(^\circ\) C for Proteinase K inactivation. After cellular lysis, 500 \(\mu\)L of tris-buffered phenol (pH 8.0) was added for extraction, tubes were mixed in vortex for 1 min and centrifuged at 12,000 \(g\) for 5 min. This procedure was repeated once.
After elimination of the organic phase (lower), 500 µL of tris-buffered (pH 8.0) phenol/chloroform/isoamyl alcohol 25:24:1 was added and tubes were mixed in a vortex for 1 min. and centrifuged at 12,000 g for 5 min. The aqueous (upper) phase was transferred to a new tube and 50 µL of 3M sodium acetate and 1 mL of ethanol PA was added, before the tubes were incubated at -20°C for 16 h for DNA precipitation. Tubes were then centrifuged at 12,000 g for 30 min, the pellet was washed with 1 mL of 70% ice cold ethanol, and the tubes were centrifuged again for 15 min at 12,000 g. Tubes were left to air dry and 200 µL of TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0) was added to resuspend the DNA; samples were stored at -20°C until analysis.

**POLYMERASE CHAIN REACTION (PCR) FOR M. pulmonis**

M. pulmonis DNA amplification was performed with 5 µL of purified samples in 25 µL reactions, containing 20 µM of each primer: sense (5’-AGCGTTTGCTTCACTTTGAA-3’) and antisense (5’-GGGCATTTCCTCCCTAAGCT-3’), 3.0 µM MgCl₂, 200 µM of each dNTP (dATP, dCTP, dGTP and dTTP) with 1.0 U of Taq DNA Polymerase in the reaction buffer supplied by the manufacturer (Invitrogen, Carlsbad, California, USA). The reaction involved an initial denaturation step for 5 min at 95°C followed by 30 amplification cycles for 1 min at 95°C, 1 min at 53°C and 1 min at 72°C, with a final extension step for 7 min at 72°C. The PCR products (266 bp) were detected by 1% agarose gel electrophoresis stained with ethidium bromide under UV light. As a negative control, H₂O was used and the UABCTIP M. pulmonis strain was used as a positive control (Institute Pasteur, Paris, France).

**EVALUATION OF THE INFLAMMATORY RESPONSE**

- Haematological analysis: platelet, red blood cell (RBCs) and white blood cell (WBCs) counts were performed in the samples obtained with EDTA, immediately after collection. Analyses were performed with a Cell Dyn 1400 device (Abbott, USA). For differential white blood cell counts, a blood smear stained by the Giemsa method was made and observed under an optical microscope.

- CRP and A1GP: serum levels of CRP and A1GP were determined in the serum samples obtained from blood collected without EDTA. They were quantified by immunoturbidimetry using commercial kits (BioTécnica, Varginha, Minas Gerais, Brazil), according to the manufacturer's instructions, using a Siemens ADVIA 1650 automated device (Siemens AG, Germany).

Statistical analysis: statistical analysis was performed using the unpaired t-test with Welsh's correction, with the aid of the software GraphPad Prism® (La Jolla, California, USA); one-tailed P values < 0.05 were considered to be significant (95% CI).

**RESULTS**

Among the studied sample, 16/20 (80%) were positive for M. pulmonis, either by culture or PCR. Samples were considered negative only with a concordance of both methods. Two samples were negative by PCR but positive by culture, and one was negative by culture and positive by PCR. Many samples presented contamination in the culture, making it impossible to identify mycoplasma growth, but were positive by PCR (13/20). Table 1 presents the CRP and A1GP results of each sample, being positive or negative for mycoplasma.

<table>
<thead>
<tr>
<th>Rat no.</th>
<th>Gender</th>
<th>Culture</th>
<th>PCR</th>
<th>CRP (mg/dL)</th>
<th>A1GP (mg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>M</td>
<td>cont</td>
<td>neg</td>
<td>4.02</td>
<td>18.19</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>cont</td>
<td>neg</td>
<td>0.62</td>
<td>9.69</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>cont</td>
<td>neg</td>
<td>3.11</td>
<td>11.93</td>
</tr>
<tr>
<td>10</td>
<td>F</td>
<td>cont</td>
<td>neg</td>
<td>1.14</td>
<td>12.97</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>pos</td>
<td>neg</td>
<td>1.49</td>
<td>7.26</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>pos</td>
<td>neg</td>
<td>3.45</td>
<td>13.87</td>
</tr>
<tr>
<td>1</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>3.99</td>
<td>6.28</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>1.69</td>
<td>9.07</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>2.07</td>
<td>14.21</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>1.51</td>
<td>8.52</td>
</tr>
<tr>
<td>11</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>4.50</td>
<td>24.58</td>
</tr>
<tr>
<td>12</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>3.85</td>
<td>18.15</td>
</tr>
<tr>
<td>13</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>4.44</td>
<td>21.82</td>
</tr>
<tr>
<td>14</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>3.68</td>
<td>20.34</td>
</tr>
<tr>
<td>15</td>
<td>F</td>
<td>cont</td>
<td>pos</td>
<td>5.00</td>
<td>24.01</td>
</tr>
<tr>
<td>16</td>
<td>M</td>
<td>neg</td>
<td>pos</td>
<td>4.21</td>
<td>29.86</td>
</tr>
<tr>
<td>17</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>5.23</td>
<td>25.51</td>
</tr>
<tr>
<td>18</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>8.20</td>
<td>25.71</td>
</tr>
<tr>
<td>19</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>12.82</td>
<td>30.68</td>
</tr>
<tr>
<td>20</td>
<td>M</td>
<td>cont</td>
<td>pos</td>
<td>5.57</td>
<td>19.51</td>
</tr>
</tbody>
</table>

Cont: contaminated culture; M: male; F: female; SP4: positive culture in SP4 medium.
CRP results were significantly higher ($P = 0.0330$), in $M. \text{ pulmonis}$ infected rats (mean = 4.48, SD = 0.70) than in those without mycoplasma infection (mean = 2.22, SD = 0.80). The same happened to A1GP ($P = 0.0321$), with a mean of 18.71 (SD = 2.005) in $M. \text{ pulmonis}$ infected rats and a mean of 13.20 (SD = 1.800) in rats without mycoplasma (Figure 1).

In the same manner, haematological tests provided additional evidence of inflammation in mycoplasma-infected animals. These results are summarised in table 2. Three samples of mycoplasma-infected rats had inadequate EDTA blood samples for haematological analysis.

It was possible to observe (Figure 2) that monocytes ($P = 0.0180$) presented higher and platelets ($P = 0.0396$) had lower levels in mycoplasma-infected rats than in those non-infected animals. For RBCs ($P = 0.0775$), Ht ($P = 0.0581$) and Hb ($P = 0.0537$) showed borderline results, demonstrating a tendency towards anaemia in the infected group, which is compatible with chronic infection. For the other parameters, there was no significant difference (Figure 2).

**Table 2** – Haematological parameters [median (SD)] in the blood samples obtained from rats with (pos) or without (neg) $M. \text{ pulmonis}$ infection

<table>
<thead>
<tr>
<th>M. pulmonis</th>
<th>RBCs mill/mm$^3$ (SD)</th>
<th>Ht % (SD)</th>
<th>Hb g/dL (SD)</th>
<th>WBCs k/mm$^3$ (SD)</th>
<th>Neutr % (SD)</th>
<th>Linfo % (SD)</th>
<th>Mono % (SD)</th>
<th>Eos % (SD)</th>
<th>Plats k/µL (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>neg</td>
<td>7.87 (0.19)</td>
<td>39.95 (0.93)</td>
<td>14.45 (0.40)</td>
<td>3.25 (0.64)</td>
<td>24 (4.4)</td>
<td>68 (5.8)</td>
<td>6 (0.4)</td>
<td>3 (1.7)</td>
<td>816 (22)</td>
</tr>
<tr>
<td>pos</td>
<td>7.27 (0.36)</td>
<td>36.54 (1.82)</td>
<td>13.15 (0.65)</td>
<td>4.14 (0.51)</td>
<td>32 (3.4)</td>
<td>59 (3.2)</td>
<td>8 (0.7)</td>
<td>2 (0.5)</td>
<td>751 (21)</td>
</tr>
<tr>
<td>$P$</td>
<td>0.0775</td>
<td>0.0581</td>
<td>0.0537</td>
<td>0.1575</td>
<td>0.1082</td>
<td>0.1417</td>
<td>0.0108</td>
<td>0.2282</td>
<td>0.0396</td>
</tr>
</tbody>
</table>

RBCs: red blood cells; Ht: haematocrit; Hb: haemoglobin; WBCs: white blood cells; Neutr: segmented neutrophils; Linfo: lymphocytes; Mono: monocytes; Eos: eosinophils; Plats: platelets.

**DISCUSSION**

As previously mentioned, mycoplasma infection in laboratory animals may cause important morbidity and mortality, especially in those used for long duration studies, or in those with a subclinical infection that may be complicated by experimental procedures. However, the most important impact might be its interference in interpreting experimental results. For instance, if a studied group presents a higher level of inflammation or immune system activation, one may not be certain whether this finding is due to the intervention being studied, or the eventual presence of mollicute infections. On the other hand, control groups, if also infected, may bring an important bias to the result.

In this direction, the aim of this study was to evaluate the level and effect of $M. \text{ pulmonis}$ infection in rats bred by the animal facility of the University of Blumenau, and discuss the impact of the use of these animals in scientific experimentation.
RBCs: red blood cells; Ht: haematocrit; Hb: haemoglobin; WBCs: white blood cells; Neutr: segmented neutrophils; Linfo: lymphocytes; Mono: monocytes; Eos: eosinophils; Plats: platelets; * P < 0.02; ** P < 0.05.

Figure 2 – Graphical representation of the haematological parameter results in the blood samples obtained from rats with (pos) or without (neg) M. pulmonis infection.
It was observed a level of 80% of *M. pulmonis* infection. Around the world, similar rates have been found as well, for instance in Taiwan, with a 40% *M. pulmonis* infection rate. In Western Europe facilities, there were better sanitary conditions than might be expected, in which rates can be as low as 3%. The level of *M. pulmonis* infection found in the animal facility of our institution makes the use of these animals in experimental models that evaluate immunity and/or infections by other microorganisms simply impossible.

With this study, we not only had information about the level of mycoplasma infection in laboratory rodents in our reality, which was not available before, but also observed a significant activation of the inflammatory response, and therefore an immune modulation, in the infected animals group. This information is not only important to researches that might be using this kind of animal for experimentation, but may also serve as an alert for researchers and those responsible for animal breeding to check the practices and quality of the facility, in case it was not available, as in ours. This is essential to assure the supply of high quality rodents for research.

This high prevalence of *M. pulmonis* in rats bred at one of the most important facilities of our State constitutes a severe issue, requiring prophylactic and preventive measures to be applied. The use of molecular methods to monitor facilities may constitute an expressive advantage, but, as it was observed, some samples may present inhibitory substances in the DNA sample, being positive only by culture, in spite of the fact that the later method might be more subject to contamination. It is also worth noticing that the inflammatory parameters evaluated are acute-phase proteins, thus, they indicate the presence of an inflammatory process in progress in respiratory tract. One should also consider that the observed results may not be exclusively due to *M. pulmonis* infection, but also to other recognized respiratory agents such as the cilia-associated respiratory bacillus, a frequent pathogen co-infecting rats along with *M. pulmonis*. Evaluations of an animal facility for experimental purposes should include screening for these organisms as well.

**CONCLUSION**

It is considered that, before developing experimentations evaluating the immune modulation or effects of other infectious diseases models, researchers must verify the conditions of the animal facility to offer the required animals, if this is not automatically provided. More than that, it would be advisable that the animal facility could ship a certificate of quality along with the animals, indicating the management type applied to breeding, and every sanitary test executed. As a further step, it was observed that an organised and directed effort for the accreditation of facilities in every country would be of great interest for the scientific community, for example, through the Association for Assessment and Accreditation of Laboratory Animal Care International, the Federation of Laboratory Animal Science Associations, among other entities.

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**CONFLICTING INTERESTS**

None declared.

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**Infecção por micoplasma e efeitos inflamatórios em roedores criados em laboratório para pesquisa experimental**

**RESUMO**

*Mycoplasma pulmonis* é o agente da micoplasmose respiratória murina, sendo a espécie mais frequentemente isolada em reprodução de laboratórios. Uma das importantes consequências de sua infecção é a redução de taxas de reprodução e significantes interferências nos experimentos usando esses animais. Sua infecção experimental é também considerada um modelo para estudos de infecção pelo *M. pneumoniae*. Atualmente foram relatadas taxas de infecção por *M. pulmonis* de até 75% em roedores criados em locais de pesquisas experimentais; no entanto, este efeito no sistema imunológico dos animais ainda não foi avaliado. Portanto, o objetivo deste estudo foi o de investigar o efeito do *M. pulmonis* em roedores criados em instalações da Universidade de Blumenau pela reação em cadeia por polimerase, a fim de verificar a viabilidade do uso desses animais para experiências. Parâmetros de inflamação tais como proteína C reativa (PCR) e alpha-1 glicoproteína ácida (AGPA), assim como os parâmetros hematológicos foram determinados por meio de amostras de sangue. Foi observada taxa de positividade de 80% de infecção por *M. pulmonis* e, também, que a infecção por micoplasma estava associada ao aumento dos níveis de PCR e AGPA, assim como os monócitos. Considerando que a possibilidade da infecção por micoplasma ser um assunto conhecido e que exames de rotina e medidas preventivas têm sido internacionalmente padronizados por muitos anos, os achados deste estudo indicam que a utilização dos animais avaliados é inaceitável para a pesquisa experimental. É extremamente aconselhável que os laboratórios que conduzem esse tipo de pesquisa estejam seguros quanto à ausência de infecção por micoplasma nos animais que foram usados.

**Palavras-chave:** Ratos; Micoplasmas; Pneumopatias; Hematologia; Química Clínica; Imunologia.
Infección por micoplasma y efectos inflamatorios en roedores criados en laboratorio para investigación experimental

RESUMEN
Mycoplasma pulmonis es el agente de la micoplasmosis respiratoria murina, siendo la especie más frecuentemente aislada en roedores de laboratorios. Una de las importantes consecuencias de la infección es la reducción de las tasas de reproducción y significativas interferencias en los experimentos usando esos animales. Su infección experimental está también considerada un modelo para estudios de infección por M. pneumoniae. Actualmente han sido relatadas tasas de infección por M. pulmonis de hasta un 75% en roedores criados en locales de investigaciones experimentales; sin embargo, este efecto en el sistema inmunológico de los animales todavía no ha sido evaluado. Por lo tanto, el objetivo de este estudio fue el de investigar el efecto del M. pulmonis en roedores criados en instalaciones de la Universidad de Blumenau por la reacción en cadena de la polimerasa, con el fin de verificar la viabilidad del uso de esos animales para experiencias. Parámetros de inflamación tales como proteína C reactiva (PCR) y alpha-1 glicoproteína ácida (AGPA), así como los parámetros hematológicos se determinaron a través de muestras de sangre. Se observó una tasa de positividad de 80% de infección por M. pulmonis y, también, que la infección por micoplasma estaba asociada al aumento de los niveles de PCR y AGPA, bien como de los monocitos. Considerando que la posibilidad de que la infección por micoplasma es un tema conocido, y que los exámenes de rutina y las medidas preventivas han sido internacionalmente estandarizadas durante años, los hallazgos de este estudio indican que la utilización de los animales evaluados es inaceptable para la investigación experimental. Se aconseja vehementemente que los laboratorios que conducen este tipo de investigación estén seguros con relación a la ausencia de infección por micoplasma en los animales que fueron usados.

Palabras clave: Ratas; Micoplasmas; Enfermedades Pulmonares; Hematología; Química Clínica; Inmunología.

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