Influence of a DNA-hsp65 vaccine on bleomycin–induced lung injury*

Influência do biofármaco DNA-hsp65 na lesão pulmonar induzida por bleomicina

Adriana Ignacio de Padua1, Célio Lopes Silva2, Simone Gusmão Ramos3, Lúcia Helena Faccioli4, José Antônio Baddini Martinez5

Abstract

Objective: To evaluate the effects of immunization with a DNA-hsp65 vaccine in an experimental model of pulmonary fibrosis. Methods: A total of 120 male C57BL/6 mice were distributed into four groups: SS, injected with saline (placebo) and then receiving intratracheal (IT) instillation of saline; SB, injected with saline (placebo) and then receiving IT instillation of bleomycin; PB, treated with plasmid, without bacterial genome, and then receiving IT instillation of bleomycin; and BB, treated with the vaccine and then receiving IT instillation of bleomycin. Bleomycin was instilled 15 days after the last immunization, and the animals were killed six weeks thereafter. The left and right lungs were removed, the former for morphological analysis and the latter for hydroxyproline measurements. Results: The proportion of deaths within the first 48 h after the IT instillation (deaths attributed to the surgical procedure) was higher in the SB group than in the SS group (57.7 vs. 11.1%). The mean area of pulmonary interstitial septa was greater in the SB and PB groups (53.1 ± 8.6% and 53.6 ± 9.3%, respectively) than in the SS and BB groups (32.9 ± 2.7% and 34.3 ± 6.1%, respectively). Conclusion: Immunization with the DNA-hsp65 vaccine reduced the deposition of noncollagen matrix in a model of bleomycin–induced lung lesion.

Keywords: Pulmonary fibrosis; Bleomycin; Collagen.

Resumo

Objetivo: Avaliar a influência do biofármaco DNA-hsp65 em um modelo de distúrbio fibrosante pulmonar experimental. Métodos: Foram estudados 120 camundongos machos C57BL/6, divididos em quatro grupos: grupo SS, animais tratados com salina (placebo) e injetados com salina intratraqueal (IT); grupo SB, tratados com salina (placebo) e injetados com salina IT; grupo PB, tratados com plasmídeo, sem gene bacteriano, e injetados com bleomicina IT; e grupo BB, tratados com DNA-hsp65 e injetados com bleomicina IT. A bleomicina foi injetada 15 dias após a última imunização, e os animais sacrificados seis semanas após o uso da droga IT. O pulmão esquerdo retirado foi utilizado para análise morfológica, e o pulmão direito para dosagens de hidroxiprolina. Resultados: A proporção de camundongos que apresentaram morte não-programada depois de 48 h da injeção IT foi maior no grupo SB em comparação ao grupo SS (57.7% vs. 11.1%). A área percentual média de interstício septal foi maior nos grupos SB e PB (53.1 ± 8.6% e 53.6 ± 9.3%, respectivamente) em comparação aos grupos SS e BB (32.9 ± 2.7% e 34.3 ± 6.1%, respectivamente). Os grupos SB, PB e BB mostraram aumentos nos valores médios da área de interstício septal corada por picrosírius em comparação ao grupo SS (SS: 2.0 ± 1.4%; SB: 6.2 ± 4.9%; PB: 7.2 ± 4.2%; e BB: 6.6 ± 4.1%). O conteúdo pulmonar de hidroxiprolina no grupo SS foi inferior ao dos demais grupos (SS: 104.9 ± 20.9 pg/pulmão; SB: 160.4 ± 47.8 pg/pulmão; PB: 170.0 ± 72.0 pg/pulmão; e BB: 162.5 ± 39.7 pg/pulmão). Conclusão: A imunização com o biofármaco DNA-hsp65 interferiu na deposição de matriz não-collagênica em um modelo de lesão pulmonar induzida por bleomicina.

Descritores: Fibrose pulmonar; Bleomicina; Collágeno.

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**Introduction**

Pulmonary interstitial diseases constitute a heterogeneous group of illnesses in which varying degrees of inflammation and fibrosis can be observed. Idiopathic pulmonary fibrosis (IPF) is a distinct type of chronic fibrosing interstitial pneumonia that is progressive, and the outcome is typically fatal. It constitutes the paradigm of interstitial pulmonary diseases, both for its frequency and for its severity. Currently, there are no truly efficacious options for treating this disease.

Among the many factors that can influence the scarring process and the final degree of fibrosis is the type of immune-inflammatory response involved. Among other factors, some studies indicate that T lymphocytes play an important role in the physiopathological process of IPF. It is known that, in the interstitium, responses involving T helper 2 (Th2) lymphocytes predominate in patients with this disease. In pulmonary lesions caused by this disease there are mastocytes and eosinophils, as well as increased quantities of interleukin (IL)-4 and IL-13. The finding of reduction of the expression of the Th1 cytokine interferon-gamma (IFN-γ), which can activate cellular mechanisms of antigen removal and restoration of healthy tissue, can have implications in the establishment of pulmonary fibrotic lesions.

A study involving lung biopsies of 10 patients with IPF characterized the types of cellular infiltrates using immunohistochemical techniques. The majority of mononuclear cells infiltrating the tissue (> 50%) were stained for IL-4 and IL-5, whereas < 7% were stained for IFN-γ. In addition, there is accumulating evidence that the macrophages of patients with pulmonary fibrosis present increased expression of IL-10 and IL-13, cytokines equally related to Th2 inflammatory responses. This set of observations suggests that the predominance of a Th2 immune-inflammatory response in pulmonary fibrosis over Th1 responses can contribute to the development of pulmonary fibrosis.

Recently, a vaccine based on the technology of the recombinant DNA, which uses the 65-kDa heat shock protein (hsp65) as the antigen, was developed in Brazil and has shown promising results in the prevention and treatment of tuberculosis in rodents. Studies have shown that, in animals infected with the tuberculosis bacillus and exposed to this antigen, the initial immunological response is altered, promoting the expression of Th1 cytokines, with a consequent increase in IFN-γ and decrease in the numbers of bacterial colony-forming units. Due to these findings, we raised the hypothesis that immunization of animals with the DNA-hsp65 vaccine promotes protection against inflammatory and fibrosing alterations induced by the intratracheal (IT) instillation of bleomycin in mice.

**Methods**

A total of 120 male C57BL/6 mice, eight to ten weeks of age and free of specific germs, were studied. Throughout the experimental period, the animals were housed on ventilated shelves, in facilities appropriate for the maintenance of germ-free animals, at the Central Animal Facility of the Department of Clinical Medicine of the Faculty de Medicina de Ribeirão Preto da Universidade de São Paulo (FMRP-USP, University of São Paulo at Ribeirão Preto School of Medicine). The study was approved by the FMRP-USP Committee for Ethics in Animal Experimentation.

The mice were randomized into four groups:

1) SS group: injected with saline (placebo) and then receiving intratracheal (IT) instillation of saline
2) SB group: injected with saline (placebo) and then receiving IT instillation of bleomycin
3) PB group: animals immunized with the DNA-hsp65 vaccine without bacterial genome—plasmid only—and then receiving IT instillation of bleomycin
4) BB group: treated with the DNA-hsp65 vaccine and then receiving IT instillation of bleomycin

**Table 1** - Number of animals and proportion of unexpected deaths in the studied groups of animals.

<table>
<thead>
<tr>
<th>Group</th>
<th>Initial number</th>
<th>Early mortality, n (%)</th>
<th>Number of surviving animals &gt; 48 h</th>
<th>Late mortality, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SS</td>
<td>26</td>
<td>8 (30,8)</td>
<td>18</td>
<td>2 (11,1)</td>
</tr>
<tr>
<td>SB</td>
<td>32</td>
<td>6 (18,8)</td>
<td>26</td>
<td>15 (57,7)*</td>
</tr>
<tr>
<td>PB</td>
<td>30</td>
<td>6 (20,0)</td>
<td>24</td>
<td>11 (45,8)</td>
</tr>
<tr>
<td>BB</td>
<td>32</td>
<td>9 (28,0)</td>
<td>23</td>
<td>6 (26,1)</td>
</tr>
</tbody>
</table>

SS: animals injected with saline (placebo) and then receiving intratracheal (IT) instillation of saline; SB: animals injected with saline (placebo) and then receiving IT instillation of bleomycin; PB: animals immunized with the DNA-hsp65 vaccine without bacterial genome—plasmid only—and then receiving IT instillation of bleomycin; and BB, treated with the DNA-hsp65 vaccine and then receiving IT instillation of bleomycin. *p < 0.05 in comparison with the SS group and through Fisher’s exact test.
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2) SB group: injected with saline (placebo) and then receiving IT instillation of bleomycin
3) PB group: animals treated with the DNA-hsp65 vaccine without bacterial genome (plasmid only), and then receiving IT instillation of bleomycin
4) BB group: animals treated with the DNA-hsp65 vaccine and then receiving IT instillation of bleomycin

The DNA-hsp65 vaccine was developed by cloning a DNA fragment of the genome that encodes the hsp65 gene of M. leprae in the pcDNA3 plasmid, as previously described. The animals were immunized with injections into the quadriceps, at a dose of 100 µg and a volume of 50 µl per dose. Corresponding volumes of plasmid only or saline were administered in the control groups. The immunization schedule encompassed four doses administered with a two-week interval between each dose.

In order to induce lung injury, bleomycin sulfate (Blenoxane®; Bristol Myers-Squibb, São Paulo, Brazil) was administered IT at 0.15 U for each 30 g of weight, two weeks after the last dose of vaccine, plasmid or saline. On this occasion, the animals were initially anesthetized with 2.5% tribromoethanol (Sigma-Aldrich, St. Louis, MO, USA) at 1 mL for each 100 g of weight, injected intraperitoneally. The animals were then immobilized with adhesive tape on a wooden plank, in the supine position and with the head hyperextended. A median cervical incision of 1-1.5 cm was made, with dilatation of the structures and exposure of the trachea. The corresponding doses of bleomycin diluted in distilled water were aspirated, and the final volume was brought up to 0.1 mL, using a syringe and a thin hypodermic needle, aiming to promote better distribution of the solution in the airways. Corresponding volumes of saline were similarly instilled in the control groups. At the end of the instillation, the cervical incision was sutured with fine cotton thread.

The animals were killed six weeks after IT instillations. Prior to sacrifice, the mice were anesthetized with intraperitoneal injections of 2.5% tribromoethanol and a cervical incision was made. After exposure, the trachea was sectioned and a small-diameter plastic catheter was introduced. An abdominal incision was made, and the descending aorta was sectioned while the animal was still alive and breathing spontaneously. The mice were then killed by exsanguination, after which the chest was opened and the cardiopulmonary block was removed for dissection. Using a fine cotton thread, the right main bronchi was ligated and sectioned. When freed, the right lung was weighed and stored at −70 °C for later biochemical analysis. Buffered 10% formalin was infused through the thin plastic catheter at a pressure of 20 cmH₂O. The extremity of the trachea was then ligated with cotton thread and the plastic catheter was removed, after which the formalin-inflated lung was fixed by immersion in a receptacle also containing formalin. Left lungs were stored in formalin for later histological studies.

After standard processing, sagittal sections of left lungs were submitted to histological examination. The fixed lungs were stained with hematoxylin-eosin (H&E) and picrosirius. The H&E-stained sections were mounted on slides and morphometrically analyzed by a pulmonary pathologist specialized in histological alterations of the septal interstitium. The findings are expressed as percentages of the total area of the field. In each slide, 10 randomly

![Figure 1](image_url) - Percentage data corresponding to the area of the septal interstitium stained with hematoxylin-eosin in the different groups: SS: animals injected with saline (placebo) and then receiving intratracheal (IT) instillation of saline; SB: animals injected with saline (placebo) and then receiving IT instillation of bleomycin; PB: animals immunized with the DNA-hsp65 vaccine without bacterial genome—plasmid only—and then receiving IT instillation of bleomycin; and BB, treated with the DNA-hsp65 vaccine and then receiving IT instillation of bleomycin. *p < 0.05 through analysis of variance and Student-Newman-Keuls post-test.
early deaths was equal among the groups, the late death rate was significantly higher in the SB group than in the SS group. Although the percentage of late deaths was lower in the BB group than in the SB and PB groups, the difference was not statistically significant.

The mean percentage of the interstitial septal matrix stained with H&E was significantly lower in the BB group than in the SB and PB groups, although it was comparable to that found for the SS group (BB: 34.3 ± 6.1%; SS: 32.9 ± 2.7%; SB: 53.1 ± 8.6%; PB: 53.6 ± 9.3%; Figure 1). The mean percentage of the interstitial septal matrix stained with picrosirius was significantly lower in the SS Group than in the other groups (SS: 2.0 ± 1.4%; SB: 8.2 ± 4.9%; PB: 7.2 ± 4.2%; BB: 6.6 ± 4.1%; Figure 2).

The total lung hydroxyproline content was significantly lower in the SS group than in the other groups (SS: 104.9 ± 20.9 pg/lung; SB: 160.4 ± 47.8 pg/lung; PB: 170.0 ± 72.0 pg/lung; BB: 162.5 ± 39.7 pg/lung; Figure 3). Figure 4 illustrates the histological aspects of the studied groups. In those images, the histological damage induced by

Results

The numbers of animals designated for each group are listed in Table 1, together with the early and late death rates. Although the percentage of
The DNA-hsp65 vaccine was initially developed in order to promote protection against infections caused by *M. tuberculosis*. However, there is mounting evidence that this vaccine, in addition to protecting animals against tuberculosis infection, can act as a therapeutic agent when the disease is already established. The vaccine was obtained by cloning the gene that encodes hsp65 in plasmids. To that end, plasmids expressing the promoter of the immediate early gene of the cytomegalovirus, or another originating from hydroxymethylglutaryl coenzyme A reductase in mice, were employed, with similar results.\(^{(15)}\)

Many studies show that the DNA-hsp65 vaccine is an immunomodulatory agent and a powerful stimulant of the cellular immune response with efficacy superior even to that of the bacillus Calmette-Guérin.\(^{(11-15,21,22)}\) Its intramuscular administration prior to *M. tuberculosis* infection leads to an elevation of the number of hsp65-reactive T cells. This increase occurs simultaneous to a drop in the numbers of CD4\(^+\)/CD8\(^−\) cells, as well as of CD4\(^−\)/CD8\(^+\) cells. Immunization with the vaccine also leads to increased numbers of splenic T cells that express the CD44-hi phenotype, a marker associated with Th1 type responses. Regardless of their phenotype, hsp65-reactive cells from immunized animals produce elevated levels of IFN-\(γ\). Conversely, the lymphocytes of unvaccinated animals respond to hsp65 stimuli in vitro, producing predominantly IL-4.

The administration of the vaccine eight weeks after inoculation with the tuberculosis bacilli, in addition to reducing the number of germs at the tissue level, also leads to similar immunological alterations, that is, deviation of a Th2 immune response to a Th1 pattern. It is of note that animals not infected with the tuberculosis bacillus and treated only with the vaccine also present a significant elevation in the number of lymphocytes expressing IFN-\(γ\) in comparison with unvaccinated control animals.\(^{(21)}\)

These findings support the application of the new agent in clinical scenarios in which increasing the cellular immune response could be beneficial. In a pilot study, the intratumoral application of the DNA-hsp65 vaccine in patients with advanced phase head or neck carcinoma was well tolerated and was, in some cases, associated with a reduction of tumor volume.\(^{(23)}\)

As previously mentioned, studies conducted principally with material from lung biopsies and
Figure 4 - Photomicrographs of representative lung histological sections of the animals in the studied groups. A and B: SS group, no significant alterations. C and D: SB group, compact area with mixed inflammatory infiltrate (lymphomonocytic and neutrophil) and collagen deposit. E and F: PB group, presence of areas of tissue repair, mostly centered in the principal bronchial axis, with extension to the adjacent parenchyma and characterized by mixed infiltrate and collagen formation. G and H: BB group: injuries restricted to the lung axis characterized by alveolar collapse, lymphomonocytic infiltrate and new tissue repair. Staining: A, C, E, F: hematoxylin-eosin; B, D, G, H: picrosirius. Magnification of all photomicrographs: ×50. SS: animals injected with saline (placebo) and then receiving intratracheal (IT) instillation of saline; SB: animals injected with saline (placebo) and then receiving IT instillation of bleomycin; PB: animals immunized with the DNA-hsp65 vaccine without bacterial genome—plasmid only—and then receiving IT instillation of bleomycin; and BB, treated with the DNA-hsp65 vaccine and then receiving IT instillation of bleomycin.
bronchoalveolar lavage fluid obtained from patients with IPF suggest that the excessive secretion of various cytokines classified as Th2 can contribute to the genesis of the disease. Conversely, experimental data show that administration of the Th1 IFN-γ cytokine can have a protective effect against the establishment of pulmonary fibrotic processes, although the results of studies involving the use of this substance in human beings with IPF have been disappointing.\textsuperscript{(24,25)}

We developed the present study based on the hypothesis that immune stimulation with the DNA-hsp65 vaccine could be protective against the establishment of pulmonary fibrosis. The experimental model used was bleomycin-induced lung injury. Although the model cited does not reproduce exactly the histological pattern of IPF, it has been widely used as a screening tool for the evaluation of the efficacy of drugs with therapeutic potential for pulmonary fibrotic processes. In addition, the doses of the DNA-hsp65 employed in this study were based on previous studies in which the immunization schedule proved effective in stimulating murine immunity and providing protection against tuberculosis infection.\textsuperscript{(22)}

Since the beginning of the project, we have observed elevated mortality rates among mice in the first hours following manipulation for the IT instillation of substances. This phenomenon occurred to the same degree in all groups and probably reflects the development of respiratory insufficiency and other complications related to the surgical manipulation of small-diameter tracheas. Therefore, we chose to analyze only the unexpected deaths of animals surviving the first 48 h. Although the proportion of unexpected deaths in the BB group was statistically similar to that seen in the other groups, it was substantially lower than that seen in the SB group (26.1\% vs. 57.7\%). From our point of view, this suggests that the DNA-hsp65 vaccine can bring some degree of protection against lung aggression induced by the IT instillation of bleomycin. Findings referring to the measurements of the septal interstitium evaluated through morphometry in H&E-stained material point in the same direction. These measurements provide a yardstick of the global involvement and of the degree of the pulmonary septal interstitium response to chemical injury. In this aspect, the area of the septal interstitium of the BB group was significantly inferior to that of the SB and PB groups and comparable to that of the SS group.

The picrosirius staining technique stains specifically collagen fibers. Likewise, the tissue values of the amino acid hydroxyproline reflect the degree of deposition of collagen in the organs. In the present study, the mean values of the digital densitometric determination of collagen fibers identified by the first method did not differ significantly among the SB, PB and BB groups. Lung hydroxyproline content showed similar behavior to the picrosirius staining, suggesting equally that the DNA-hsp65 vaccine, in the present model, had no effect on the degree of bleomycin-induced collagen deposition.

Since no significant differences were detected in the measurements that reflect the degree of collagen deposition, the alterations in septal thickness detected in the H&E stainings are likely explained by effects on other histological elements. The interstitial septum is composed of cellular and extracellular components. In the subjective histological evaluation of the slides by a pulmonary pathologist, the SS group presented no abnormalities, although there was remodeling of the lung architecture in the SB and PB groups. Those last two groups presented mixed cell inflammatory infiltrate, predominantly lymphoplasmacytic, and areas of tissue repair located preferentially around the bronchovascular axis, extending into the adjacent parenchyma. In relation to the BB group, less intense injuries were observed, with new tissue repair, restricted to the central axis of the airways. Therefore, the effects of the DNA-hsp65 vaccine seem to have been more evident in the reduction of the intensity of the mixed cell inflammatory infiltrates than in the process of collagen deposition per se. It is equally relevant that, in addition to collagen, the extracellular matrix is composed of other biochemical elements, such as elastin, proteoglycans, laminin and fibronectin. Therefore, the protective effects of the vaccine might have occurred in some or all these last components.\textsuperscript{(26)}

Recent advances in the understanding of the pathogenesis of IPF call into question the role of inflammatory alterations as primordial factors for the establishment of pulmonary fibrosis and suggest that epithelial elements associated with abnormal cicatriziation processes are major primordial elements of the disease.\textsuperscript{(27)} In the animal model of bleomycin-induced pulmonary fibrosis, however, previous studies indicate
the existence of a correlation between the degree of the acute inflammatory alterations and later establishment of fibrosis. In the present study, all animals were studied at the same time, that is, eight weeks after the end of immunization with the vaccine and six weeks after the IT instillation of bleomycin. We cannot exclude the possibility that sacrificing the animals at a later date, with greater evolution of the fibrotic process, might produce different results, particularly regarding a greater influence of immunization on the establishment of the fibrotic injuries.

We highlight that this is only a preliminary experimental study, and that other schedules with higher doses of the DNA-hsp65 vaccine or shorter intervals of administration might, in theory, provide better results. In addition, indeed, there are no animal models that fully reproduce the usual interstitial pneumonia-type injuries, which are the anatomopathological substrates of IPF. Therefore, the true role of the DNA-hsp65 vaccine in the treatment of pulmonary fibrosis, particularly IPF, in theory, would only be truly determined through the use of well-designed clinical assays.

In summary, we can conclude that the data here obtained indicate that immunization with the DNA-hsp65 vaccine had a positive effect on the degree of deposition of the noncollagen matrix in a model of bleomycin-induced lung injury. Additional studies are necessary in order to elucidate more fully the effects that the vaccine has on pulmonary fibrotic processes.

References


