Natural history of experimental arthritis induced by *Paracoccidioides brasiliensis* in wistar rats

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**Abstract**

**Introduction**: Paracoccidioidomycosis (PCM) is the most prevalent systemic mycosis in Latin America. This study aimed to evaluate the natural history of *Paracoccidioides brasiliensis*-induced experimental arthritis of the knee joints in Wistar rats.  

**Methods**: Rats were randomly allocated to either an absolute control group, or 15-day, 45-day, or 90-day experimental (fungus-inoculated) groups.  

**Results**: Experimental groups developed classic signs of articular PCM. Titers of anti-gp43 were observed to increase during the interval from 15 to 45 days post-inoculation.  

**Conclusions**: Articular arthritic lesions were induced and progressed during the study period in all experimental groups.  

**Keywords**: Arthritis. Paracoccidioidomycosis. *Paracoccidioides brasiliensis*.

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Paracoccidioidomycosis (PCM) is a systemic mycosis caused by the fungus *Paracoccidioides brasiliensis* (Pb), initially known as Brazilian blastomycosis, South American blastomycosis, or Lutz-Almeida-Splendore disease[1].  

PCM represents a significant public health challenge, given the susceptibility of young adults in their most productive phase of life (between 30 and 59 years), and the difficult-to-treat nature of the infection[2].  

The route of infection is generally through the upper airways, through which Pb conidia are inhaled. These infectious propagules initially settle in the lungs, but the fungus can then spread throughout the body, causing lesions in the internal mucocutaneous and osteoarticular organs[3,4].  

Osteoarticular involvement becomes chronic in about 60% of cases and occurs secondary to systemic involvement[5]. Articular PCM manifestation involves signs of intense inflammation, including functional impairment of the joint. X-ray examinations demonstrate cartilage destruction, space reduction, and joint effusion[6].  

However, few studies report evidence-based details regarding articular PCM; the understanding of its evolution is currently based on assumptions found in literature. Thus, this study aimed to evaluate the evolution of Pb-induced experimental arthritis of the knee joint in Wistar rats.  

After approval was received from the Research Ethics Committee of UNIOESTE (N. 05/2013), the study was performed in the Laboratory of Microbiology at the State University of Western Paraná, in Cascavel/PR. We used 45-60 day-old Wistar rats (n = 24) randomly and allocated them evenly to either an absolute control group (ACG, which remained uninfected) or 15-day, 45-day, or 90-day experimental (fungus-inoculated) groups (which were inoculated with Pb). The animals in the experimental groups were sacrificed after their respective experimental periods had elapsed, and ACG animals were sacrificed at the 90-day mark.

Induction of experimental arthritis took place as described by Loth et al.[7]. The animals were anesthesitized with ketamine and xylazine, intraperitoneally (50 mg/kg and 10 mg/kg body weight). Each animal received a medial knee region inoculation using a Pb18 strain cell suspension (1×10⁵ yeast cells/ml in...
anti-gp43 antibody titers as described by Ramos et al. 8. immunological analysis, including ELISA-based assessment of and sacrificed through the guillotine. Blood was collected for ketamine and xylazine (50 mg / kg, 20 mg / kg body weight) corresponding animals were anesthetized using intraperitoneal measurement was done just prior to sacrifice.

At the end of each experimental group interval, the corresponding animals were anesthetized using intraperitoneal ketamine and xylazine (50 mg / kg, 20 mg / kg body weight) and sacrificed through the guillotine. Blood was collected for immunological analysis, including ELISA-based assessment of anti-gp43 antibody titers as described by Ramos et al. 8.

For histological analysis, the right knee joint of each animal was dissected and fixed in 10% formalin buffered solution, followed by a routine protocol to embed the tissue in Paraffin. Longitudinal microsections of 7 μm thickness were stained with hematoxylin and eosin for morphological analysis, and with Grocott’s stain to evaluate Pb18 infection, prior to examination using light microscopy.

Differences between groups of ELISA absorbance readings were statistically compared using an ANOVA. Edema measurements were statistically compared using the Wilcoxon test. Both comparisons were performed using GraphPad Prism®, version 3.0 for Windows XP, Microsoft Office®, and results were considered statistically different at a 5% significance level.

Joint edema increased in the Pb18-infected group, peaking on day 15, followed by a decrease up to day 45, and then a plateau until day 90. The increase in joint edema varied between 30% and 40% of baseline thickness (Figure 1).

Titers of anti-gp43 antibodies against Pb exhibited a progressive increase until day 45, after which the level at day 90 was reduced, but still higher than that observed at day 15 (Figure 2).

Qualitative histological analysis demonstrated that Pb18-mediated synovitis was established by day 15, including granulomatous inflammation exhibiting poorly organized granulomas and edema in the perimeniscal region synovial membrane, inflammatory cells in the intima and sub-intima, and increased intimal thickness as well as neovascularization. Figure 3A shows the part of the ACG joint, with anatomical aspects preserved for comparison. Morphology of other articular tissues (cartilage of the femur and tibia), as well as of the subchondral bone was normal (Figure 3B).

At 45 days post-infection, the experimental group exhibited exacerbation of pathology, including pannus formation. Synovial membrane inflammatory infiltrates became larger, more intense, and more diffuse, affecting its full thickness and with the presence of marked neovascularization. Numerous, dense Pb18-containing granulomas were observed, whereas Langhan’s giant cells and empty Pb18 capsules were less frequently observed (Figures 3C and 3D).

At 90 days post-infection, large granulomas had formed in the synovial membrane (Figure 3E), with Pb18 localizing to joint-adjacent regions, including the perimuscular region, with areas of lytic necrosis (Figure 3F). The main pathological finding at this timepoint was the presence of Pb18 in the subchondral bone of the femur and tibia (Figure 3G and 3H).

Edema formation peaked at day 15, had somewhat abated by day 45, and had stabilized by day 90. These results agree with the findings of a similar study, in which a considerable increase in edema was also observed in the experimental groups at days 15 and 45 post-induction of infection 9.

The day 45 experimental group exhibited an increase in titers of anti-gp43 antibody against Pb18, indicating persistence and progression of the disease. However, at day 90, a reduction in antibody titers was observed. This agrees with the findings of a similar study, in which researchers monitored anti-gp43 expression at days 15 and 45 in the same experimental model, and observed an increase in titers at day 45, possibly due to fungal proliferation 10.

In this study, although dissemination and spread of the disease had occurred by day 90 as demonstrated by the
anatomical pathology examination, the animals’ immune systems failed to maintain antibody titers. This agrees with the findings of a study in which patient plasma levels of anti-gp43 antibody exhibited a significant elevation on day 28 post-infection, but a reduction on day 56 post-infection, even in the absence of treatment.11

During the evaluation of anatomical pathology, classical signs of PCM were observed in the experimental groups. At day 15 post-infection, synovitis with granulomatous inflammation, edema, an inflammatory infiltrate, and neovascularization predominated. This agrees with human case reports in which the presence of numerous Pb elements of varying sizes (including multiple budding elements) and the presence of fungus-containing Langhan’s giant cells amidst macrophages was observed.12

In this study, infection severity was higher on day 45 compared to day 15, as evidenced by pannus formation, increased synovial membrane edema, severe synovitis, severe neovascularization, periosteal granuloma, necrosis, and periarticular dissemination. Similarly, a clinical study using magnetic resonance imaging also found soft tissue reactive edema and synovial inflammation in PCM patients with joint involvement.13

During prior development of the experimental model employed in this study, intense inflammatory signs and focal necrosis occurred at day 45 of exposure.7 In the current study, lytic necrosis was only observed to be present by day 90. This may be due to virulence of the fungal strain used for inoculation; although both studies used the Pb18 strain, inoculum virulence was not verified in either study. Clinical studies, too, report the presence of well-defined osteolytic lesions in patients.14

In the current study, the presence of Pb18 was observed in joint-proximal regions, including perimuscularly, by day 90 post-infection.

Similarly, by day 90 post-infection, Pb appeared in the subchondral bone of the femur and tibia (figures 3G and 3H), underlying the entry point of the middle genicular artery. This suggests that dissemination of the fungus to this location occurred via this vessel. In a retrospective study examining seven clinical cases of osteoarticular PCM, involvement of the bony tissue was observed in all patients, with over 40% of samples exhibiting multiple bone lesions.14

The current study pioneers investigation into the evolution and later stages of experimental Pb-induced arthritis, and an attempt has been made to contextualize findings despite the paucity of prior literature. The experimental model of Pb-induced arthritis employed by this study performed satisfactorily for the purpose of studying arthritis evolution.

Based on these results, it would appear – at least in a rodent model - that untreated PCM progresses over time, including development of characteristic anatomical pathology findings such as necrosis and dissemination to adjacent tissues. Despite disease progression, however, a decrease in titers of anti-gp43 antibodies against Pb can be expected.

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REFERENCES


