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## VIRULENCE PROFILE OF TEN *Paracoccidioides brasiliensis* ISOLATES. ASSOCIATION WITH MORPHOLOGIC AND GENETIC PATTERNS

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### SUMMARY

Ten isolates of *Paracoccidioides brasiliensis* were examined for differences in virulence in outbred mice intravenously inoculated with the fungus, associated with mycelial morphology, and genetic patterns measured by random amplified polymorphic DNA (RAPD). Virulence was evaluated by viable yeast cell recovery from lungs and demonstration of histopathologic lesions in different organs. The results showed that the isolates presented four virulence degrees: high virulence, intermediate, low and non-virulence. RAPD clustered the isolates studied in two main groups with 56% of genetic similarity. Strains with low virulence, Pb265 or the non-virulent, Pb192, showed glabrous/cerebriform morphology and high genetic similarity (98.7%) when compared to the other isolates studied. The same was observed with Bt79 and Bt83 that shared 96% genetic similarity, cottony colonies and high virulence. The RAPD technique could only discriminate *P. brasiliensis* isolates according to glabrous/cerebriform or cottony colonies, and also high from low virulence strains. Isolates with intermediate virulence such as Pb18, Pb18B6, Bt32 and Bt56 showed variability in their similarity coefficient suggesting that RAPD was able to detect genetic variability in this fungal specie. Virulence profile of *P. brasiliensis* demonstrated that both mycelial morphologic extreme phenotypes may be associated with fungal virulence and their *in vitro* subculture time. Thus, RAPD technique analysis employed in association with virulence, morphologic and immunologic aspects might prove adequate to detect differences between *P. brasiliensis* isolates.

**KEYWORDS:** Antibodies; Colony morphology; *Paracoccidioides brasiliensis*; RAPD; Virulence.

### INTRODUCTION

*Paracoccidioides brasiliensis* the etiologic agent of paracoccidioidomycosis (PCM), is a thermally dimorphic pathogenic fungus. This mycosis is endemic in South America occurring from Central America to Argentina<sup>33</sup>. The different clinical forms of the disease, and the occurrence of paracoccidioidomycosis-infection, may be due to host related factors as well as to characteristics of the infecting agent, mainly virulence<sup>8</sup>.

Animal models have been very useful for studying host-parasite interactions in PCM, and have revealed that different *P. brasiliensis* isolates may vary in virulence<sup>4,12,27</sup>, a fact that might explain the different clinical manifestation of the disease.

Differences in virulence of *P. brasiliensis* isolates in mice have also been reported<sup>4,14,27</sup>, and associated with the amount of  $\alpha$ -1,3 glucan in the fungal cell wall<sup>24</sup>. Virulence is also influenced by *in vitro* passage and storage of the isolates<sup>4,22</sup>.

Some studies employed to analyze growth curves, and morphology of yeasts have shown variability between *P. brasiliensis* strains, but no association with pathogenicity has been observed<sup>12,13</sup>. Another aspect

that has been studied in connection with virulence is mycelial growth morphology. Cottonous morphology appears to be related to virulent strain while glabrous morphology is associated with low virulence strains<sup>25</sup>.

Random Amplified Polymorphic DNA (RAPD) analysis has also become very useful for microbial strain identification, and has been successfully used to detect genomic variations between fungal isolates such as *Aspergillus fumigatus*<sup>1,19</sup>, *Histoplasma capsulatum*<sup>35</sup> and *P. brasiliensis*<sup>31</sup>.

RAPD analysis of *P. brasiliensis* has been used for clustering strains by geographic area<sup>5,11</sup>, as a means to clarify PCM pathogenesis<sup>26</sup> and also, to correlate RAPD pattern with virulence<sup>18</sup>. Taking these facts together, we investigated virulence profile in ten *P. brasiliensis* isolates using different parameters and exploring the relationship between their phenotypes and RAPD pattern.

### MATERIAL AND METHODS

***P. brasiliensis* isolates:** Ten isolates of *P. brasiliensis* were employed, eight obtained from patients with paracoccidioidomycosis, one isolated from a hamster inoculated with armadillo visceral organ

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homogenates (Pb-T) and another recovered from C57Bl/6 mice infected with Pb18 (Pb18B6) (Table 1). All isolates were maintained on 2% Glucose, 1% Peptone and 0.5% Yeast Extract medium (GPY medium) at 35 °C from time of isolation and were sub-cultured to fresh media every 15 days.

**Colony morphology:** Colony morphology of the ten isolates was evaluated by observation of giant colonies in their mycelial form. Each isolate obtained from a 6 day-old culture in GPY medium at 35 °C was plated on the central area of three 220 x 100 mm Petri dishes with Potato dextrose agar (PDA) with aid of a platinum loop that permits an inoculation around 3 mm in diameter. Plates were incubated at 25 – 28 °C for eight weeks, and colonies were examined for shape, size, texture and presence of pigment.

**Animals:** Groups of 10-week-old male outbred swiss mice were provided from the animal facility of Botucatu Campus, São Paulo State University, for this study. The animals were provided with acidified water and sterilized pellet food *ad libitum*.

**Virulence studies:** Groups of 10 mice each were inoculated by the intravenous (iv) route with  $2 \times 10^6$  yeast cells from a 6 day-old culture of *P. brasiliensis* isolates: Pb18, Pb18B6, Bt32, Bt56, Bt79, Bt83, Bt84, Pb192, Pb265 and PbT. Fungus viability was determined by phase microscopy<sup>30</sup> and it was always above 90%. Four weeks post-infection animals were sacrificed and submitted to histopathologic study, recovery of viable fungi from lungs, and measurement of specific antibody levels.

**Sacrifice and histopathology:** Each animal was sacrificed by ether anesthesia and cardiac puncture exsanguination. Blood was collected and tissue samples were taken from lungs, liver, spleen, kidney, and adrenal gland. Blood samples were allowed to clot and the supernatant sera centrifuged and stored at -20 °C. Tissue samples were fixed in formalin, and embedded in paraffin, cut into 4 µm sections and stained with hematoxylin-eosin.

**Recovery of viable yeast from lungs:** The number of viable fungi in infected mouse lungs was determined by counting colony forming units (CFUs). At sacrifice, the right lungs from all animals were aseptically removed weighed and individually homogenized in 1.5 mL of sterile phosphate buffer saline pH 7.2 (PBS) in a tissue grinder. Aliquots of 100 µL of each homogenate were plated by triplicate onto plates with brain heart infusion (BHI) agar (Difco Laboratories, Detroit, Mich.) supplemented with 4% (v/v) normal horse serum and 5% *P. brasiliensis* 192 culture filtrate, the latter constituting the source of growth-promoting factors<sup>28</sup>. Plates were incubated at 35 °C and colonies were counted daily until there were no further increases. Results were expressed as logarithmic values of CFU/g of lung.

***P. brasiliensis* antigen:** Culture filtrate antigen (PbAg) was prepared using a pool of 6-day-old yeast cells from the ten *P. brasiliensis* isolates, cultured in Casal's medium<sup>7</sup> at 35 °C. Each culture filtrate was centrifuged separately at 10,000 g for 30 min at 4 °C, concentrated to a 10 mL-volume and dialyzed against PBS. Then the ten culture filtrate were pooled, filtered through a 0.22 µm Millipore membrane, and submitted to sterility tests. Protein concentration was determined by the method of LOWRY *et al.*<sup>16</sup>.

**Humoral immunity:** Specific antibody levels to *P. brasiliensis* were measured by the enzyme linked immunosorbent assay (ELISA) as described previously<sup>17</sup>. All sera were tested in two-fold dilutions and each PbAg was used at 20 µg protein/mL. Goat-anti-mouse total immunoglobulin peroxidase conjugate (Sigma, Chemical Co, St Louis, Mo, USA) was used in all experiments. As controls, antibody titers of sera from non-infected mice, housed under identical conditions as the infected animals, were also determined. Average optical density obtained with control mice sera at 1:16 dilution was considered the reaction cut off value. Optical densities for each dilution of the experimental sera were compared with control values. Sample titers were expressed as the reciprocal of the highest serum dilution with absorbance higher than cut off value.

**DNA-preparation:** DNA was prepared by PAN & COLE's<sup>21</sup> method to obtain protoplast cell. After cell wall digestion, protoplast was resuspended in buffer lyses, 50 mM Tris-HCL (Sigma) and 20 mM EDTA pH 7.5 (Sigma), 250 µL SDS 10%, 300 µL Proteinase K (10 mg/mL) (Sigma) and incubated at 65 °C for 30 min. Before DNA purification, protoplast cells buffer lyses were incubated on ice with 5M potassium acetate for 60 min, and then treated with RNase. DNA was then purified with phenol, phenol:chlorophorm, chlorophorm, isopropanol, and 70% ethanol; DNA was then resuspended in 10 mM Tris-HCl plus 1 mM pH8 EDTA and stored at -20 °C.

**RAPD analysis:** Seventeen 10-bp primers (Operon Biotechnology) namely, OPB05, OPB07, OPB8, OPB10, OPB11, OPB12, OPB14, OPB15, OPB17, OPB18, OPG13, OPP02, OPP14, OPr13, OPr09, OPR08, and OPR07 were used for the RAPD assay<sup>10,34</sup>. The DNA amplification reactions were performed in a 13 µL system that consisted of 5 ng genomic *P. brasiliensis* DNA; 1.3 µL of 10x Taq buffer (200 mM Tris-HCl; pH 8.4; 500 mM KCl); 2.5 mM of each deoxynucleotide triphosphate; 3 µL of primer (5 ng/µL); IU Taq DNA polymerase (Gibco, BRL, Geithersburg, MD, USA); 1.5 mM of MgCl<sub>2</sub>; 1.04 µL non-acetylated bovine serum albumin (10 mg/mL) (New England Biolabs, Beverly, MA, USA) and water to make up the volume to 13 µL. The reaction underwent 40 cycles in an automatic thermal cycler (MJ Research PT 100 Mini-Cycler); each cycle consisting of 1 min at 92 °C for denaturation, 1 min at 35 °C for annealing, 2 min at 72 °C for elongation. Randomly amplified products underwent electrophoresis analysis on a 1.4% agarose gel and were developed with ethidium bromide staining.

**Statistical analysis:** The RAPD pattern from each isolate was analyzed by NTSYS 1.7 software (Numerical Taxonomy and Multivariate Analysis System, NY, USA), using the Jaccard coefficient<sup>23</sup>. The dendrogram was derived from the distance matrix by the Unweighted Pair-Group Method Arithmetic Average (UPGMA). Number of CFU/g of lung was transformed into logarithmic units and analyzed by one-way analysis of variance (ANOVA) followed by multiple comparisons by the Tukey test. Antibody level from sera was analyzed by the Kruskal-Wallis test. Correlation between CFU/g of lung and antibody levels was done by linear correlation. Significance was considered at  $p < 0.05$ .

## RESULTS

**Colony morphology:** Phenotypes were distinguished by differences

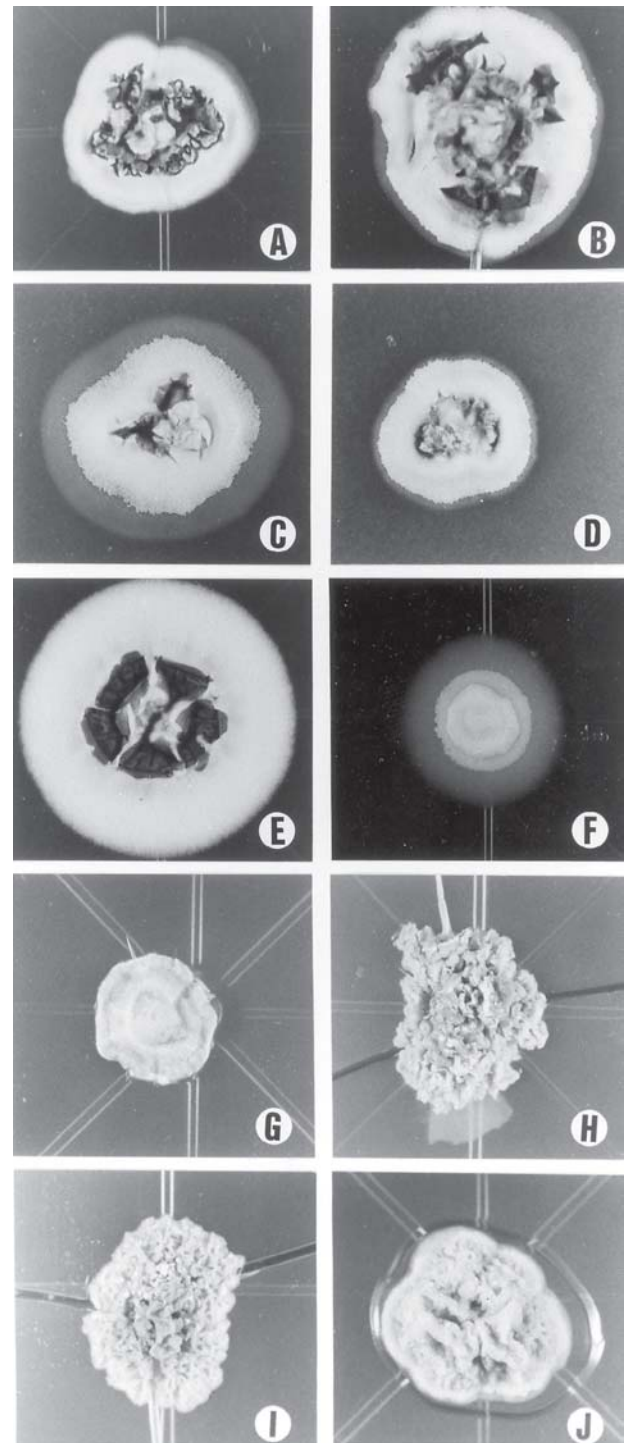
in colony shape, texture, and presence of pigment. The colonies of *P. brasiliensis* isolates presented two distinct phenotypes which could be maintained and stabilized in subculture. The first was present in Pb18, Pb18B6, Bt32, Bt79, Bt83 and Bt84 isolates, all of which showed cottony phenotype with colonies adherent to the medium, white cotton-like surface with or without central fissures and the undersides were brownish in color. In the second phenotype, colony color was creamy, wrinkled and folded, aerial hyphae were rare, and the colonies never became floccose or cottony. This type was named glabrous phenotype and included Bt56, Pb192, Pb265 and PbT isolates (Fig. 1).

**Recovery of viable *P. brasiliensis* from lungs:** Yeast cells counts (CFUs) in the lungs of mice inoculated with the different *P. brasiliensis* isolates revealed that the infection varied according to the isolate employed. The ten isolates showed four different pathogenic degrees of virulence according to number of viable fungal units recovered from infected mouse lung tissue: a) high virulence was attributed to Bt79, Bt83, and Bt84, with significantly higher CFU values than all the other isolates; b) intermediate virulence to Bt56, Bt32, Pb18, and Pb18B6; c) low virulence to Pb265 and PbT; and d) non-virulence to Pb192 only (Table 1). There was no difference in the virulence profile between Pb18 isolate sub-cultured long term *in vitro* in our laboratory and Pb18B6 originating from Pb18 recovered from C57Bl/6 infected mice.

**Histopathological studies:** Differences in virulence could also be observed in the histopathologic analysis. Seven of the ten isolates studied induced lung lesions. Animals infected with Pb18B6, Bt32 and Bt56 had few lung lesions that covered less than 1/4 of the histological section (Fig. 2 A). Mice inoculated with Bt84 and PbT had moderate lesions with 1/4 to 1/2 of the histological section involvement and those with Pb18, Pb192 and Pb265 isolates did not have lung lesions. Mice infected with Bt79 and Bt83, classified as highly virulence isolates, presented extensive lung lesions that covered 2/3 or the whole section (Fig. 2B). Hepatic lesions were only detected in animals infected with Bt79, Bt83, Bt84, and Bt56. In Bt56 infected mice, there were few hepatic lesions and no fungi were seen. Extensive spleen and adrenal gland lesions were only observed in Bt79 and Bt83 infected animals (Fig. 2C) with high virulence. Low or non-virulent isolates did not induce lesions upon infection.

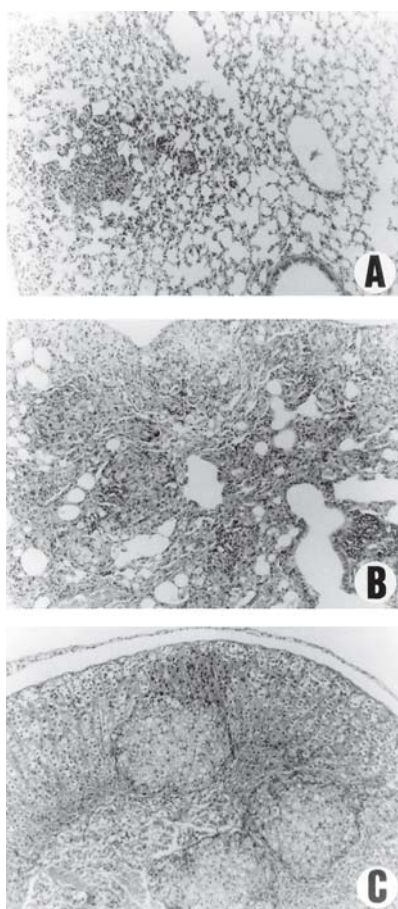
**Antibody levels:** *P. brasiliensis* isolates elicited distinct serological profiles during infection (Table 1). Mice infected with Bt79, Bt83 and Bt84 had antibody titers more elevated in comparison to those inoculated with the other seven isolates, and significantly higher than those detected in mice infected with Pb192. In these animals antibodies were absent or very low, with titers ranging from 0 to 1. Animals infected with intermediate and low virulence isolates had similar antibody levels (Table 1).

**RAPD analysis:** A total of 103 reproducible amplification products were sufficiently polymorphic to allow strain differentiation. Depending on primer, one to seven bands were separated, ranging in size from 0.25 to 4.5 kb. RAPD analysis showed two groups with 56% genomic identity, group one clustered Bt32, Pb192, Pb265, Pb18, PbT, Bt79, Bt83, Pb18B6 strains and group two Bt56 and Bt84 (Fig. 3). Group one was divided in two subgroups, the first had all isolates except Pb18B6 and the second subgroup corresponded to Pb18B6 alone. Genetic similarity between these two subgroups was about 88%. First



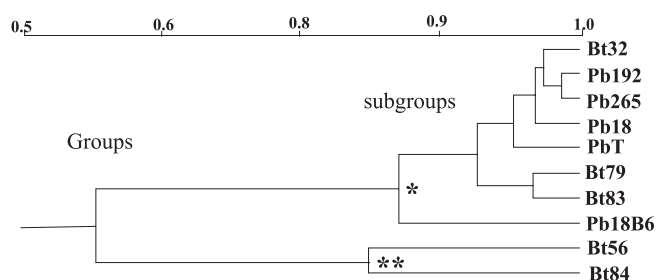
**Fig.1** - Giant colonies from *Paracoccidioides brasiliensis* isolates cultured at 25 °C on Potato dextrose agar medium. A) Pb18, B) Pb18B6, C) Bt79, D) Bt83, E) Bt84, F) Bt32, G) Bt56, H) Pb192, I) Pb265, J) PbT.

subgroup isolates shared more than 90% of genetic similarity. Isolates Pb265 and Pb192 had a similarity coefficient of 98.7% and the mycelia colony form were glabrous/cerebriform with little difference in virulence pattern. Similarity of 96% was seen between Bt79 and Bt83,



**Fig. 2** - Lesion histopathology: (A) Lung tissue from mice infected with Bt32. (B) Lung tissue and (C) adrenal gland from mice infected with Bt79 isolate. (HE - 100 x).

and mycelium morphology was similar for both isolates. Bt79 and Bt83 had cottony mycelia morphology with black central fissure and high virulence, demonstrated by extensive lesions in lungs, liver and adrenal gland of the infected mice.



**Fig. 3** - Dendrogram of the relationship between *Paracoccidioides brasiliensis* isolates based on UPGMA clustering of matrix obtained by Jaccard coefficient. Note presence of two main groups: group 1 (\*) and group 2 (\*\*)

## DISCUSSION

In this study we report the presence of phenotypic variations in *P. brasiliensis* isolates according to colonial morphology. The fungus was able to adopt either a glabrous or a cotton-like appearance. Many organisms have acquired the ability to vary their phenotype, generating variants within infectious populations as a mechanism designed to escape threatening environmental changes, such as host immunity development<sup>32</sup>. From the human isolates studied here, cotton-like colonies were more frequent than glabrous, as previously seen in other studies of the morphologic and physiologic features of *P. brasiliensis* isolates<sup>25</sup>.

Virulence of the ten fungal isolates as determined by both, number of viable fungi recovered from infected mouse lung tissues and histopathologic analysis were shown to be associated suggesting their efficiency in determining virulence of *P. brasiliensis* isolates. According to SINGER VERMES *et al.*<sup>29</sup> estimation of the number of viable fungi in infected mouse organs by CFUs count is one of the most direct and trustworthy methods to assess severity of the disease. Fungal isolates Bt79, Bt83 and Bt84 which presented the cottony phenotype were classified as highly virulent strains as based on the high number of yeast cells, recovered from the lungs (5.04 to 6.10 log CFUs/g), and also, on the isolates capacity to disseminate to extrapulmonary organs (liver and spleen), leading to extensive granulomatous lesions. Thus, the degree of virulence was inferred from disease severity in the mice.

**Table 1**  
Comparative characteristics and degree of infection between 10 *Paracoccidioides brasiliensis* isolates

Isolate	Morphology	LogCFU/g lung tissue (mean ± SD)	Virulence	Antibody titer	Subculturing time (years)
Pb 18	Cottony	2.05 ± 0.41*	Intermediate	48 (8 - 128)	> 30
Pb 18/B6	Cottony	2.60 ± 0.46 *	Intermediate	48 (8 - 128)	3
Bt 32	Cottony	2.47 ± 0.50 *	Intermediate	12 (4 - 64)	11
Bt 56	Glabrous	3.12 ± 0.89 *	Intermediate	64 (2 - 128)	6
Bt 79	Cottony	6.10 ± 0.91 #	High	2560 (256 - 4096) *	2
Bt 83	Cottony	5.34 ± 0.7 #	High	256 (128 - 256)*	2
Bt 84	Cottony	5.04 ± 0.62 #	High	256 (64 - 512)*	2
Pb 192	Glabrous	0.0	Avirulent	1 (0 - 16)	> 30
Pb 265	Glabrous	1.05 ± 0.37♦	Low	128 (16 - 256)	> 30
Pb T	Glabrous	1.07 ± 0.38 ♦	Low	128 (128 - 128)	5

Antibody titer: results are expressed as median values with the range in parentheses; \*(p < 0.05) versus Pb192 (Kruskal-Wallis test). Log CFU/g lung tissue: \*(p < 0.05) versus PbT, Pb265 and Pb192; #(p < 0.01) versus Bt56, Bt32, Pb18, Pb18B6, PbT, Pb265 and Pb192; ♦(p < 0.05) versus Pb192 (ANOVA)

On the other hand, low virulent Pb265, PbT, and non-virulent Pb192 isolates, having glabrous morphology, were recovered from lungs in low numbers (less than 1.07 log CFUs/g) or not at all and, additionally, did not induce histopathologic lesions upon inoculation. These results suggest that both extreme phenotypes may be associated with fungal virulence, and, also, in connection with their record of *in vitro* sub-culturing. In general, fungi with glabrous morphology had been maintained by longer sub-culturing periods (5 to 30 years) than cottony ones (2 to 3 years). The exception occurred with Pb18 and Bt32 isolates, which presented intermediate virulence, lower fungal recovery (2.47 log CFUs/g or less) and cottony phenotype, maintained after long subculturing time (11 to 30 years). Thus, our results are in accordance to those obtained by SANO *et al.*<sup>25</sup>, showing that *P. brasiliensis* isolates with low *in vitro* subculture time and cottony morphology frequently present high virulence, while isolates with cottony phenotype, but long *in vitro* subculture period present lower or intermediate virulence. The phenomenon of *in vitro* virulence attenuation has also been described for other fungi that cause deep mycosis, such as *Blastomyces dermatitidis*<sup>3</sup>, and *Cryptococcus neoformans*<sup>9</sup>. Data from *in vivo* studies have revealed that virulence of *P. brasiliensis* isolates could be attenuated or even lost after *in vitro* long-term sub-cultivation<sup>4,14,22</sup>. This phenomenon could be reverted by re-isolation of the fungus from mice, independent of their susceptibility or resistance to paracoccidioidomycosis<sup>14</sup>.

Serum levels of specific antibodies to *P. brasiliensis* were significantly higher in animals infected with the highly virulent isolates and might well be associated with the antigenic fungal load in mouse tissues. These results appear similar to those in human disease, where high antibody levels are not associated with protection but with progressive fungal dissemination to various host organs<sup>6,8</sup>.

RAPD analysis of the ten *P. brasiliensis* isolates showed two main groups with 56% genomic identity, suggesting the variability of this fungus. Our results showed that *P. brasiliensis* recovered from patients from the same city or state were highly variable in contrast with the results obtained by CALCAGNO *et al.*<sup>5</sup> who found correlation between RAPD patterns and geographical region from which each fungal strain was isolated. Further studies on our RAPD grouping could be helpful ascertaining classification and fungal variability.

RAPD profile could discriminate some *P. brasiliensis* Bt pairs, such as Bt79 and Bt83 highly virulent with a similarity coefficient of about 96%, and also Pb192 and Pb265 with 98.7% genetic similarity and low virulence. Each of these pairs had similar virulence degree in infected mice, similar colony morphology in PDA medium, with mice showing similar antibody titers. Our results suggest an association between RAPD patterns and degree of virulence in *P. brasiliensis* as previously suggested<sup>18</sup>. However, the RAPD technique did not discriminate between intermediate virulence strains such as Pb18, Pb18B6, Bt32 and Bt56 which showed variability in their similarity coefficient. Also, Pb18 strain and Pb18B6, the isolate derived from Pb18 when recently isolated from severely infected C57Bl/6 which exhibited only 88% of genetic similarity and the same capacities of infecting mice. These results are not in accordance with MOTTA *et al.*<sup>20</sup> who detected 100% similarity between Pb18 and its isolate derivatives (virulent, attenuated, and non-virulent) with different capacities of infecting susceptible mice. Thus, RAPD analyses could not discriminate strain virulence from one large group, but only between some pairs of *P. brasiliensis* strains in the subgroups studied.

In this study Pb18 and Pb18B6 showed no differences in virulence and morphology parameters, but some genetic variability. The genetic differences between Pb18 and its derivative Pb18B6 might be attributed to changes in the genomic DNA of the fungus required to adapt to the host's environment. Genetic changes or microevolution on standard strains have been suggested for other fungi such as *C. neoformans*<sup>9</sup>, *B. dermatitidis*<sup>15</sup> and *H. capsulatum*<sup>2</sup>, and have been associated with phenotypic changes and loss of virulence.

Together, the results presented here demonstrate that *P. brasiliensis* isolates may vary in colony morphology and virulence associated with *in vitro* sub-culturing time. RAPD analysis showed some correlation with virulence and morphology of *P. brasiliensis* isolates. This technique allows detection of genetic variations between Brazilian isolates of *P. brasiliensis*<sup>5,11,31</sup>. Differences in virulence between *P. brasiliensis* isolates may be dependent on factors that allow fungal growth under adverse conditions in the host. Elucidation of the factor which allows his fungus to overcome host defenses will lead to a better understanding of the pathogenesis of paracoccidioidomycosis.

## RESUMO

### Perfil de virulência de dez isolados de *Paracoccidioides brasiliensis*. Associação com morfologia e padrão genético

Dez isolados de *P. brasiliensis* foram avaliados em relação à patogenicidade por inoculação intravenosa em camundongos e associação com morfologia miceliana e padrão genético por amplificação genômica do DNA polimórfico (RAPD). A patogenicidade, avaliada por recuperação de fungos viáveis a partir de tecido pulmonar e por lesões histopatológicas em diferentes órgãos, mostrou que os isolados apresentaram quatro graus de virulência: alta virulência, virulência intermediária, baixa virulência e não virulência. A técnica de RAPD agrupou os isolados em dois grupos com 56% de similaridade genética. Amostras com baixa virulência Pb265 ou não virulência Pb192 apresentaram morfologia glabra/cerebriforme e alta similaridade genética (98,7%) quando comparadas com os outros isolados estudados. O mesmo foi observado com os isolados Bt79 e Bt83, que compartilharam 96% de semelhança genética, colônias cotonosas e alta virulência. Essa técnica pode discriminar apenas isolados com morfologia glabra da cotonosa e com alta e baixa virulência. Isolados com virulência intermediária como Pb18, Pb18B6, Bt32 e Bt54 mostraram variabilidade no coeficiente de similaridade, sugerindo que a técnica de RAPD permite mostrar variabilidade genética nessa espécie fúngica. O estudo do perfil de virulência das amostras de *P. brasiliensis* demonstrou que os dois fenótipos extremos de morfologia miceliana podem ser associados com a virulência do fungo e com o tempo de subcultivo *in vitro*. Assim, a análise de RAPD, utilizada em conjunto com aspectos de virulência, morfológicos e imunológicos pode ser considerada adequada para detectar diferenças entre isolados de *P. brasiliensis*.

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