EXPERIMENTALLY INDUCED INVASIVE ASPERGILLOSIS IN MICE

ABSTRACT: In this study systemic response to intravenous administration of *Aspergillus fumigatus* conidia was investigated. The intensity of response was evaluated by a survival rate and by histopathological tissue analysis. Administration of all doses (10^6 — 5x10^7) of *Aspergillus fumigatus* conidia caused mortality, but the highest mortality and the shorter time of survival were noted at higher doses applied. At the highest applied dose, the presence of spores and hyphae was noted in lungs and kidneys. Histological analysis revealed the presence of intense inflammatory reaction in lungs, kidneys and spleen. Functional and histological changes observed provide means to study both mechanisms and drug interventions in systemic *Aspergillus* infection.

KEY WORDS: *Aspergillus fumigatus*, mice, systemic aspergillosis, urinary obstruction

INTRODUCTION

Species of genus *Aspergillus* are representatives of saprophytic filamentous fungi found in most environments. The most common species of *Aspergillus* causing invasive diseases include *A. fumigatus*, *A. flavus*, *A. niger*, *A. clavatus*, *A. glaucans*, *A. nidulans*, *A. terreus* and *A. versicolor*. *Aspergillus* sp. is weak pathogen, but might cause a disease in an immunocompromised host. Aspergillosis comprises a variety of infection, manifestation including invasive aspergillosis, pulmonary aspergilloma and allergic bronchopulmonary aspergillosis (Denning, 1998). Diseases that cause *A. fumigatus* are very difficult to diagnose, because the most diagnostic features are not specific, and patients are usually asymptomatic.

Animal experimental models offer an approach for *Aspergillus* infection studies. By using animal individuals, with defenses impaired by glucocorticoids, cyclophosphamide etc. (immunosuppressed animals), valuable informa-
tion concerning pathogenesis of *Aspergillus* infections was obtained. Studies on immunocompetent hosts, on the other hand enable investigations of the mechanisms of resistance to *Aspergillus*.

In this study we investigated a response to systemically applied *A. fumigatus* conidia. With this aim, survival and histopathology of distinct organs were analyzed, following the intravenous injection of conidia.

**MATERIALS AND METHODS**

Conventionally housed female C57BL/6 mice, eight to twelve weeks old, are used in the experiments. Animals were housed under constant conditions (temperature 19—21°C, daily-night rhythm 12 h), with food and water *ad libitum*. The experiments were conducted with adherence to Ethical Committee of Institute for Biological Research “Siniša Stanković”.

*A. fumigatus*, human isolate, from the Institute of public health of Serbia “Dr Milan Jovanović Batut” was subcultured on standard mycological slants (Booth, 1971). Inoculum was prepared by flooding the surface of agar slants with sterile 0,85% NaCl with 0,1% Tween 80. The suspension of spores was prepared in apyrogenic sterile physiological saline and doses of 1x10⁶, 1x10⁷ and 5x10⁷ conidia were applied intravenously into each mouse. The control mice received saline solely.

Animals were inspected two times a day. All mice were observed for a total 14 days after the infection. Mice that survived until the day 14, were euthanized.

Presence of fungi in organism of mice was established by histological analyses. Tissue specimens were fixed in 4% formalin (pH 6, 9). Fixed material was dehydrated in graded ethanol series. Material is then embedded in paraffin at 57°C. Sections 5 μm thick, were stained with hematoxylin-eosin (H & E).

Specific gravity, protein and haemoglobin content in urine were determined by test strips Combur¹⁰ Test®M (Roche Diagnostics GmbH, Germany) as parameters of renal function.

Results were statistically processed by Mann-Whitney U test. As significant was considered p < 0,05.

**RESULTS AND DISCUSSION**

All doses of applied conidia induced mortality in experimental animals. Mortality rate was proportional to the injected inoculum. Inoculation of 1x10⁶ and 1x10⁷ conidia per mouse caused mortality in 40 and 60% of mice respectively. Dose of 5x10⁷ conidia caused mortality in all treated animals, by day five following inoculation (Figure 1). Higher doses also reduced time of survival (the higher dose, the shorter time of survival). Mortality/survival data are in accordance with the study showing high mortality at the dose of 10⁷ of infected immunocompetent mice (Cenci et al., 1997).
Presence of infection was assessed by observation of animal prostration. Pronounced hypodynamic state and piloerection were noted in infected mice, in accordance with the data from the studies of invasive aspergillosis in mice (Duong et al., 1998). Mice in hypodynamic state died shortly after these signs appeared.

Histological data revealed the presence of inflammatory pulmonary response (presence of lymphocytes in peribronchial and/or perivascular sites) in all experimental groups. Microabscesses were noted in lungs, liver, kidneys and spleen of animals. In animals which received the highest dose the presence of conidia and hyphae in lungs and kidneys was noted.

Renal aspergillosis, at highest dose was accompanied with compromised renal function as judged by the changes in selected urinary parameters (Figure 2). In all control individuals, specific gravity values were 1,015 (test strips values ranged from 1,000 to 1,030). In 62% of treated mice, this value was 1,030, and in the rest value of 1,025 was detected. Difference between the control and treated groups is statistically significant (p = 0,0066). Significantly increased haemoglobinuria (p = 0,023), and a tendency of increase in urine protein content (p = 0,089), were noted in treated individuals. Renal dysfunction observed in these animals, is in agreement with the studies which showed that kidneys are the primary target organs for intravenous A. fumigatus infection (Latege, 1999) in animals and humans, and with reports which de-
monstiration urinary obstruction as a consequence of renal infection detected in humans (De Medeiros, 1999; Bisi, 2003).

CONCLUSION

In conclusion, presented data demonstrated both functional and histological changes in organs of mice following systemic application of *A. fumigatus*. This model might provide means to study mechanisms of invasive aspergillosis, as well as drug testing in preclinical trials.

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LITERATURE


ЕКСПЕРИМЕНТАЛНО ИНДУКОВАНА АСПЕРГИЛОЗА КОД МИШЕВА

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Резиме

Aspergillus fumigatus je опорунтистичка гљива која се може наћи у свим срединама. Опportunистичке гљиве прорукују појаву болести код имунокомпромитованих особа, односно код особа са ослабљеним имунским системом. Најчешће болести које изазива A. fumigatus су: плућна аспергилоза, аспергилом и алергиска бронхопулмонарна аспергилоза. Ове болести су тешке за дијагнозу јер су симптоми неспецифични.

Интрахосписка апликација конидија Aspergillus fumigatus изазива промене у преживљавању и понашању животиња. Са повећањем концентрације конидија смањује се време преживљавања. Највећа апликована доза (5x107 конидија) доводи до угнчања свих животиња до петог дана након инокулације. Инфициране животиње карактеришу промене у понашању (акинезија, атонија) и пилоерекција.

Хистолошком анализом је показано присуство интензивне запаљење реакције у плућима, бубрезима и слезима третираних јединки, као и присуство спора и хифа у бубрезима и плућима јединки које су примиле највећу дозу A. Fumigatus. Код ових јединки запажен је и поремећа бubreзне функције на основу промена у специфичној тежини урина, pH вредности, присуству протеинурије и хемоглобинурије.