THE OCCURRENCE OF ASPERGILLOSIS IN FLOCK OF TURKEY POULTS

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Abstract: Aspergillosis is frequent fungal disease of different avian and mammal species, caused by infection by the fungi of genus Aspergillus. The disease is characterized by inflammatory changes in the respiratory system and sometimes has generalized onset when more organ systems are affected. In this paper, we examined a flock of turkey poults, 21 days old, at one farm in Serbia. Clinical signs of central nervous system in the form of ataxia, torticollis, paresis and paralysis of legs and wings were observed. The mortality rate in the flock was 7.2 %. In ten out of twelve necropsied turkey poults multiple yellowish-white granulomas, one to three millimeters in diameter on lungs were found. In nine out of twelve necropsied turkey poults solitary yellowish-white granuloma, three to five millimeters in diameter on sagittal section of the cerebrum or cerebellum were found. Mycological finding revealed fungi Aspergillus fumigatus. For the evaluation of histopathological changes in lung and brain and demonstration of fungal hyphae, three stain methods were used: haematoxylin-eosin (HE), Grocott methenamine silver and periodic acid Schiff (PAS) method. Microscopic examination of lung and brain has revealed the presence of granulomatous foci and caseous necrosis with surrounding region of proliferation including giant cells, macrophages, heterophils and lymphocytes and outer capsule of connective tissue. The fungal hyphae were hardly or not visible in HE stained sections, while septed and arborized hyphae were easily demonstrated by Grocott and PAS stain predominantly in central parts of granuloma. For diagnostic of mycotic infection is necessary to use different histochemical methods for evaluation of histopathological changes and detection of etiological agent, including isolation to obtain etiological diagnosis.

Key words: turkey poults, mycotic granuloma, Aspergillus fumigatus
Introduction

Aspergillosis is frequent, economically important, fungal disease of different avian and mammal species caused by fungi of the genus Aspergillus. The disease is characterized by inflammatory changes in the respiratory system and sometimes has generalized onset when more organ systems are affected. Clinical signs of aspergillosis depend upon which organ or system organ are involved and whether infection is localized or generalized. Pathomorphological substrate of this disease is mycotic granuloma (aspergiloma) which is commonly found in lungs and air sacs. Numerous reports have described encephalitic or meningoencephalitic aspergillosis in turkey poults, ducklings and chickens (Knežević and Matejić, 1996; Ivetić et al., 2003; Özmen and Dorrestein, 2004; Singh et al., 2009, Kureljušić et al., 2011a). Olias et al. (2010) have described a case of articular aspergillosis of hip joints in turkeys. Comparative experimental study of the susceptibility of different species of poultry to infection with Aspergillus fumigatus has shown that turkeys are more susceptible than chickens. Primarily, there are two forms of aspergillosis in poultry: acute and chronic. Acute aspergillosis is usually characterized by severe outbreaks in young birds and high morbidity and high mortality. The mortality rate may range from 70 to 90%. Chronic aspergillosis occurs in adult breeder birds (particularly turkeys). It is very difficult to diagnosticate this infection by clinical examination because in many cases more organ systems are affected. It is known from literature that the diagnosis of fungal infections of poultry often be placed only after histopathological examination of organs of infected poultry (Richard et al., 1984, Saif, 2008). For diagnostic of mycotic infection is necessary to use more methods including clinical examination of flock, macroscopic examination of carcasses, isolation of pathogens, as well as histochemical methods for detection tissue lesions and fungal elements. Histochemical methods which can be used are routine hematoxylin-eosin staining method, periodic acid-Schiff (PAS) and specific staining method for fungi Grocott methenamine silver method (GMS) (Özmen and Dorrestein, 2004). There is also a possibility to use immunohistochemical methods for direct in situ evidence of etiological agent (Jensen et al., 1997).

Materials and methods

In this paper, we examined the carcasses of 12 turkey poults, three weeks old which originated from one farm in Serbia. The mortality rate in the flock was 7,2%. Clinical examination of the flock revealed moderate depression of many turkey poults and nervous symptoms which were expressed in the form of ataxia, torticollis, paresis and paralysis of legs and wings. Food consumption was reduced leading to reduced weight gain. After necropsy, for histopathological and
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Mycological examination samples of lung and brain tissue were taken. Samples for histopathological analyses were fixed in 10% buffered formalin, routinely processed and embedded in paraffin blocks. Paraffin sections about 5 μm were stained with hematoxylin-eosin (HE), periodic acid Schiff (PAS) and Grocott methenamine silver (GMS) methods. Samples for mycological examination were inoculated onto Sabouraud dextrose agar and incubated at temperature of 250°C under aerobic conditions for isolation of infectious agents. To avoid bacterial contamination in the substrate was added 20 IU/ml penicillin G and 40 mg/ml streptomycin sulfate. Macroscopic and microscopic examination of colonies was performed according to Quinn (2002). Samples of feed and litter were also taken for mycological analysis.

**Results and discussion**

External examination of turkey poults carcasses indicates less feathering of the skin especially in the region of chest.

In ten out of twelve necropsied turkey poults granulomatous pneumonia with multiple yellowish-white granulomas, one to three millimeters in diameter were found (Figure 1).

![Figure 1. Lung of turkey poult, granulomatous pneumonia](image)

In addition, multifocal granulomatous airsaculitis of thoracic and abdominal air sacs was also observed. In advanced cases of aspergillosis, the organism can sporulate on the surface of the caseous lesions and on the walls of the thickened air sacs, as evidenced by visible greenish-gray mold growth. That was also found in one case of aspergillosis in black swan (Kureljušić et al., 2011b), but in this case there were no such lesions. In 3 out of twelve necropsied turkey poults solitary
yellowish-white granuloma of irregular shape, three to five millimeters in diameter on sagittal section of the cerebrum or cerebellum were found (Figure 2).

In two out of twelve cases granulomas were found on the cranial part of the cerebrum, and in one case granuloma was on the central part of cerebellum. Data from literature indicate that granulomas caused by Aspergillus fumigatus (aspergilomas) usually occur in the lungs and air sacs although there are data on the occurrence of aspergilomas in the cerebellum, only in turkey poult (Schulz, 1991; Ozmen and Dorrestein, 2004; Saif 2008, Kureljušić et al, 2011a). Similar to our observations, the granulomas occur either in cerebrum or the cerebellum (Saif, 2008). Rarely, aspergillomas occur simultaneously in both parts of brain (Saif, 2008). In addition to this finding, in turkey poult that were necropsied, hyperemia of cerebral blood vessels was found. In other organs there were no macroscopically visible changes.

Mycological examination revealed that the ethiological agent is fungus Aspergillus fumigatus. Growth in the medium was observed after 24 hours, and after four days, grown colonies were white and about 2 cm in diameter. By the seventh day the colony diameter increased to 3.5 cm, and there was a color change in the central part of the colonies from blue-green to the gray-green, while the edges remained white (Figure 3). Based on macroscopic and microscopic characteristics of colonies, the culture was identified as Aspergillus fumigatus.
For the evaluation of histopathological changes and demonstration of fungal hyphae, three stain methods were used: haematoxylin-eosin (HE), Grocott methenamine silver and PAS method. Histological finding in the lung and in the brain was the same. Granulomatous reactions with central caseous necrosis were observed in the HE stained slides. There was surrounding region of proliferation including giant cells, macrophages, heterophils and lymphocytes and outer capsule of connective tissue (Figure 4, Figure 5).

The fungal hyphae were either hardly visible or not visible in HE slides, while septed and arborized hyphae were easily demonstrated in all samples by
Grocott methenamine silver and PAS stain, predominantly in central parts of granuloma (Figure 6, Figure 7).

In some cases, in the sections of brain and lung stained by HE method, granulomatous changes were characterized only by presence of smaller or larger number of necrotic foci, which are infiltrated with macrophages, giant cells and heterophil granulocytes. These changes are described in the literature as early initial lesions (Jensen et al., 1997). In three out of twelve cases in HE stained sections, fungal hyphae were not apparent. Grocott and PAS methods in all twelve cases revealed septed and arborized hyphae. Thus, the specific staining methods revealed fungal hyphae even if minimal histopathological changes occurs. Based on this finding it can be assumed that the occurrence of lesions in the lung and brain depends not only on the presence of infectious agents, but also on the reaction of the immune system of the host. PAS and Grocott method are described in the literature as a very reliable method for direct visualization of the different fungal elements in tissues (Ozmen and Dorrestein, 2004; Singh et al., 2009).

Mycological analysys of litter revealed Aspergillus fumigatus while samples of feed were negative for this microorganism. It is known from literature that contaminated litter is often the source of Aspergillus fumigatus (Saif 2008).

**Conclusion**

Microscopic finding of typical mycotic granulomas in the lung and brain with central caseous necrosis and fungal hyphae surrounded by macrophages, giant cells, lymphocytes, heterophils and capsule of connective tissue pointed to aspergillosis. For diagnostic of mycotic infection is necessary to use more methods including clinical examination of flock, macroscopic examination of carcasses,
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isolation of pathogens, as well as histochemical methods for detection of tissue lesions and fungal elements.

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Pojava aspergiloze u jatu ćurica

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Rezime

Aspergiloza je često gljivično oboljenje različitih vrsta ptica i sisara, izazvana gljivicama roda Aspergillus. Oboljenje se karakteriše inflamatornim promenama u respiratornom sistemu, a ponekad može da se javi u generalizovanoj formi kada je zahvaćeno više organskih sistema. U ovom radu, ispitano je jato ćurica, starosti 21 dan, na jednoj farmi u Srbiji. Kliničkim pregledom jata ustanovljeni su simptomi oboljenja centralnog nervnog sistema u formi ataksije, tortikolisa, pareze i paralize nogu i krila. Stopa mortaliteta u jatu je bila 7,2 %. Kod 10 od 12 obdukovanih ćurica na plućima su ustanovljeni multipli žuto-beli granulomi, od jedan do tri milimetara u prečniku. Kod 9 od 12 obdukovanih ćurica na sagitalnom preseku malog ili velikog mozga nađeni su pojedinačni žuto-beli granulomi, veličine od tri do pet milimetara u prečniku. Mikološkim ispitivanjem izolovana je gljivica Aspergillus fumigatus. Za procenu histopatoloških promena u plućima i mozgu kao i dokaz hifa gljivica, korишene su tri histohemijske metode: hematoksilin-eozin (HE), Grocott methenamine silver (GMS) i PAS metoda. Mikroskopskim pregledom uzoraka pluća i mozga bojnih HE metodom, ustanovljeno je prisustvo granuloma sa centralnim područjem kazeozne nekroze koje je bilo okruženo sa gigantskim celijama, makrofagima, heterofilima, limfocitima i spoljašnjom kapsulom sačinjenom od vezivnog tkiva. Hife gljivica, kao sastavni delovi granuloma, bile su teško uočljive, a u nekim uzorcima bojnim HE metodom nisu se ni videle. Nasuprot ovome, u preparatima bojnim Grocott i PAS metodom jasno su se isticale septirane i razgranate hife u centralnim delovima granuloma.

U dijagnostici ovog gljivičnog oboljenja neophodno je primeniti više histohemijskih metoda kako za procenu histopatoloških promena tako i za dokaz uzročnika, uključujući i izolaciju u cilju postavljanja etiološke dijagnoze.
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