HISTOPLASMOSIS IN ANIMALS

Introduction

Histoplasmosis was the most common endemic mycosis in dogs in the United States and Canada between 1964 and 1976 (1). The case rates per 100,000 patient-years-at-risk at 14 colleges of veterinary medicine for histoplasmosis was 2.5 times higher than blastomycosis and 3.5 times higher than coccidioidomycosis (1). The relative risk in dogs today is unknown. In humans in the United States, histoplasmosis was the most common cause for hospitalization, 1.5 times more common than coccidioidomycosis and 4.4 times more common than blastomycosis (2). Many veterinarians are unaware that histoplasmosis is the most common endemic mycosis.

Histoplasmosis usually causes pulmonary and/or disseminated disease. While histoplasmosis is not transmissible from animal to human, concurrent infection is not uncommon because of shared exposure. Familiarity with the clinical manifestations may alert a veterinarian to consider the diagnosis. Antigen detection in urine and serum may provide a rapid diagnosis, precluding the need for invasive procedures to obtain specimens for organism identification in many patients. Antibody testing may be useful in cases with negative results by antigen testing. Itraconazole is the treatment of choice, and therapy may be monitored by antigen testing. Itraconazole absorption and metabolism vary considerably, at times causing undetectable or toxic blood levels, and blood level measurement is encouraged to assure adequate drug exposure.

Epidemiology

Histoplasmosis caused by Histoplasma capsulatum variety capsulatum is endemic in certain parts of North and South America. In some endemic areas, histoplasmosis was the most common systemic mycosis in animals. Between 1964 and 1976, 14 schools of veterinary medicine in the United States and Canada participated in a study of systemic mycoses and noted rates per hundred thousand patient years of 62 for histoplasmosis, 25 for blastomycosis and 17 for coccidioidomycosis (1).

In a study of necropsy findings in dogs from Kentucky, 36.1% had positive cultures for H. capsulatum compared to 0.7% for Blastomyces dermatitidis (3). In a similar study from Loudoun County, Virginia H. capsulatum was isolated from 44% of apparently healthy dogs and cats (4). In Kentucky, 47% of dogs (5) and 50% of Thoroughbred horses exhibited Histoplasma skin test reactivity, while only 7.3% of horses demonstrated Blastomyces skin test reactivity (6). Histoplasmosis was twice as frequent in animals from rural areas as from urban areas (3). Histoplasmosis also occurs outside of the traditional endemic area (7;8).

Several dog breeds have been shown to have an increased risk of histoplasmosis, including the Pointer, Weimaraner and Brittany spaniel (2). Mean age at diagnosis from one large study was 3.6 years (2). Cats appear to have a similar incidence of histoplasmosis as that seen in dogs. Persian cats are slightly over-represented, while Siamese cats are marginally under-represented (55). Interestingly, indoor-only cats remain at risk for
histoplasmosis (49, 88). Cases also occur in horses (9-20), lamas (21), sea mammals (22-24) and wild animals (8;25-36)

Pathogenesis

Histoplasmosis is caused by inhalation of microconidia or hyphal fragments. Although intestinal lesions are prominent in dogs with disseminated histoplasmosis, experimental infection by gastric inoculation failed to induce disease in dogs (37). All mammals are susceptible to histoplasmosis, but cases have been reported most often in dogs, cats, and horses. Birds, because of their higher body temperature, are not susceptible to natural infection (38) but may be infected experimentally, causing infection localized to their feathers (39). Histoplasmosis is not transmissible from patient to patient, or from animals to man, but disease in an animal may be a harbinger of infection in humans who were exposed at the same time (40).

Cellular immunity is critical in defense against H. capsulatum, based on analysis of risk factors for severe disease. The microconidia are inhaled and in the lungs they attract dendritic cells, neutrophils and macrophages, which phagocytose the organism, which transform into yeasts and multiply unchecked in the non-immune subject. During the first two weeks, the infection progresses and disseminates hematogenously throughout the reticuloendothelial system. By day 14 of infection, specific T cell immunity develops, halting proliferation of the yeast and progression of the infection. Evidence for self-limited dissemination includes demonstration of calcified granulomas in the spleen and liver in healthy individuals in endemic areas for histoplasmosis, which contain non-viable organisms and occasional isolation of H. capsulatum from extrapulmonary specimens in patients with acute pulmonary histoplasmosis.

Cytokines that are most important in immunity to H. capsulatum include IL-12, IL-18, TNF-α and interferon-γ. A successful T cell response requires dendritic cells, CD4 and CD8 T lymphocytes and activated macrophages. T cells produce interferon-γ and tumor necrosis factor-α, which activate macrophages to kill Histoplasma yeast. If these elements are impaired, the yeast. The importance of TNF-α in humans is highlighted by the emerging recognition of histoplasmosis as a major opportunistic infection in patients treated with TNF inhibitors.

While histoplasmosis is self-limited in over 95% of healthy humans, chronic and/or progressive disease may occur more often in animals. Thirty one to 44% of euthanized dogs and cats in endemic areas had evidence for histoplasmosis (41-43). Chronic infection also is common in bats (44;45). Interestingly the tissue reaction in bats was minimal or absent, possibly explaining their inability to eradicate the organism (44-47). The infection rate varied markedly in different genera of bats, suggesting genetic differences in susceptibility (48). Histoplasma was not isolated from wild-caught mice, suggesting that their immune response was able to kill the organism (38).

Clinical Presentation

The severity of clinical manifestations correlates with the intensity of exposure and the underlying health of the exposed individual. Cole described rapidly progressive fatal course over two to four weeks in 10% of dogs with histoplasmosis, and chronic progressive course over two to 20 months in 90% (49). Demonstration of positive cultures of pulmonary and extrapulmonary tissues of apparently healthy dogs and cats from endemic areas
suggest that the clinical findings may be overlooked in many cases. Syndromes most commonly identified include pneumonia, mediastinal lymphadenitis, and progressive disseminated histoplasmosis.

**Pulmonary.** Pneumonia is the most common manifestation in humans, and probably in animals. Dogs usually present with signs of fever, dyspnea, cough and lethargy. Radiographic findings characteristically include diffuse nodules, referred to as "cotton tuft" lesions (50) (Figure 3) or diffuse interstitial infiltrates, often accompanied by hilar lymphadenopathy (51). Alveolar infiltrates are rarely seen (51).

**Mediastinal Lymphadenitis.** Enlarged hilar or mediastinal lymph nodes may impinge upon the airways and cause cough and respiratory distress (52;53). Radiographs show tracheobronchial lymphadenopathy usually accompanied by interstitial pneumonia. The outcome has ranged from spontaneous resolution to progressive obstruction of the airways and death. Concurrent dissemination may occur (53).

**Progressive Disseminated Histoplasmosis.** Fever, weight loss, reduced activity, anemia, and interstitial lung disease are the most common manifestations in cats (54), while diarrhea, intestinal blood loss, anemia and reduced activity predominate in dogs (49;55) (Table 1), but any tissue may be involved.

Bone lesions are common in cats (54). Central nervous system and ocular lesions may be found in all animal species. Endocarditis also has been reported, noted in seven of 17 (41%) necropsy cases in dogs (57). Other tissues commonly involved at necropsy include liver, spleen, abdominal lymph nodes, and less frequently adrenal glands, kidneys, and pancreas (49).

Pulmonary involvement occurs in most cases and is usually manifested as labored breathing (54;55). Radiographs typically show diffuse interstitial, miliary or nodular infiltrates (54).

Abnormal physical findings include hepatomegaly, splenomegaly, eye lesions or discharge, subcutaneous nodules, and skin lesions (54;55). The common laboratory abnormalities are anemia, leukopenia, thrombocytopenia, hypoalbuminemia, increased liver enzyme activity, creatinine elevation, and hypercalcemia (58). The untreated course ranges from subclinical chronic infection to a rapidly fatal illness.

**Equine Abortion.** Infections in the fetus or neonatal foal may occur, causing the mare to abort or the foal to die soon after birth (18;20;59). Pulmonary and disseminated involvement usually are present in the fetus or newborn (20). In most cases the mare appears healthy but the placenta is involved.

<table>
<thead>
<tr>
<th>Table 1. Clinical findings in dogs (55) and cats (54;56)</th>
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<tr>
<td>Finding</td>
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<td>Fever</td>
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<td>Weight Loss</td>
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<td>Respiratory Symptoms</td>
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<td>Chest Radiograph Abnormal</td>
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<tr>
<td>Lethargy/Depression</td>
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<tr>
<td>Intestinal-Diarrhea, Bleeding</td>
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<td>Hepatomegaly</td>
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<td>Splenomegaly</td>
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<td>Lymphadenopathy</td>
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<tr>
<td>Eye Lesions or Discharge</td>
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<tr>
<td>Bone Lesions</td>
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<tr>
<td>Skin Lesions</td>
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<td>Anemia</td>
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<td>Hepatic Enzyme ↑</td>
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*NS = not stated
Diagnosis

Prompt diagnosis offers the greatest chance for recovery from histoplasmosis, made possible by early therapy (60). Today most cases are diagnosed by detection of Histoplasma antigen in the urine and/or serum or demonstration of yeast in the body fluids or tissues. Antibody detection may be useful in cases in which antigen tests and/or pathology are negative or specimens are not available for pathology.

Pathology. As most reports required demonstration of Histoplasma yeasts in tissues or body fluids or positive culture for diagnosis (54;55), the sensitivity of pathology is uncertain. In humans the detection of antigen in the urine or serum is more sensitive than pathology or culture (61). Other limitations of pathology are the requirements to perform invasive procedures to obtain specimens for evaluation and that the pathologists are experienced with recognition of fungal organisms.

Antigen Detection. A galactomannan antigen in the cell wall of proliferating Histoplasma yeasts is released into the tissues and blood, and excreted in the urine. Antigen was detected in the urine of 94% of cats (82) and dogs [Cunningham in press] with histoplasmosis. The highest sensitivity is achieved by testing both urine and serum. Antigen also may be detected in the respiratory secretions in patients with pulmonary histoplasmosis and cerebrospinal fluid of those with meningitis. Antigen levels decline during treatment and increase with relapse (62), providing a tool for monitoring therapy.

The antigen found in histoplasmosis cross reacts with that found in blastomycosis (63). Furthermore, the clinical findings and endemic distribution overlap. Thus, differentiation of the two mycoses may be difficult, but treatment is the same, reducing the need to distinguish histoplasmosis and blastomycosis. Antibody testing may distinguish these two mycoses (64).

Culture. Culture is rarely performed, but usually is positive in disseminated cases (42). The major limitation is the slow growth rate and risk to laboratory personnel from exposure to the mycelial form of the fungus.

Antibody Detection. Antibody detection has not been adequately evaluated in animals with histoplasmosis. The sensitivity of complement fixation (CF) was reported to be about 90% (65). Others reported a sensitivity of CF to be only 11% (57). The CF test is often uninterpretable in dogs because their serum is anti-complementary, and CF is not offered commercially at veterinary reference laboratories. Agar gel immunodiffusion (AGID) is offered commercially, but the sensitivity was only 25% in one report (3). Positive results have been reported in cats (50) and horses with histoplasmosis (20). Sensitivity may be improved using an enzyme immunoassay, which has been described in blastomycosis (63) and under development for histoplasmosis (66).

Molecular Techniques. PCR has been reported to be positive in the tissues of several animal species with histoplasmosis, mostly in dogs (67-71) but also in other species (22;23;29;67;68;72). Most reports, however, describe results in single cases. No studies have reported the sensitivity and specificity and none have reported results on body fluids, or compared PCR to other diagnostic methods. Additional studies are needed to assess the role of PCR for diagnosis of histoplasmosis.
Treatment

Guidelines for treatment are provided, but textbooks and other reviews should be used for more thorough instruction on antifungal treatment in animals. Most information is based on experience in cats treated with fluconazole or itraconazole (73) but amphotericin B is preferred for more severe cases. Relapse has occurred after stopping therapy in 40% of successfully treated cases (73).

Itraconazole. The usual dosage is 5 (74) to 10 mg/kg (75;76) given once or twice daily. At least 4-6 months of therapy is typically administered. A study in cats found itraconazole to be effective in six of 13 cases, while 3 were euthanized or died, four required a change in therapy to fluconazole because of lack of response or toxicity (73). Four of the 13 cats experienced a relapse. Treatment should be continued until at least 1-2 months after resolution of clinical signs and probably until antigen is no longer detected in urine, based on experience in blastomycosis (62). Antigen concentration should be monitored at 3 and 6 months after discontinuation of therapy, and upon recurrence of clinical findings suggestive for relapse. In another report of five cases treated with itraconazole for 3 to 8 months, all responded to therapy and remained well with follow-up from 24 to 70 months (77). Amphotericin B was given for two treatments in 2 of the 5 cases (77).

Brand-name itraconazole (Sporanox®, Janssen Pharmaceuticals) or generic itraconazole should be used, as compounded powder formulations have poor bioavailability (78) (23). Blood levels should be measured 14 (dogs) to 21 (cats) days after beginning therapy, and the preferred range is 3.0 to 10.0 µg/mL as measured by bioassay, and at least 1.0 µg/mL by HPLC. Blood levels above 10.0 µg/mL may cause more toxicity and are unnecessary for response to therapy: dosage may be reduced in patients with blood levels above 10.0 µg/mL. Inability to achieve concentrations above 3 µg/ml or toxicity are reasons to change to fluconazole. Itraconazole is eliminated by hepatic metabolism through cytochrome P450 3A4, and blood levels may be affected by medications that interact with that enzyme.

Itraconazole may cause a variety of adverse effects, most commonly loss of appetite, anorexia, vomiting, or diarrhea, which may be related to high blood levels (79). Bilirubin and hepatic enzymes also may be elevated, in association with clinical evidence for hepatitis in some cases; and should be monitored during therapy. Serum alanine aminotransferase (ALT) greater than 200 U/L may warrant discontinuation of itraconazole (80). Itraconazole may be restarted at half of the former dose. Ulcerative dermatitis was also observed in 7.5% of dogs receiving itraconazole at 10 mg/kg/d (81).

Fluconazole. Response to fluconazole was similar to that with itraconazole in cats (73). Of 17 cats treated with fluconazole, 3 died or were euthanized, 9 completed therapy, one switched to itraconazole because of poor response, and 4 relapsed. Some veterinarians prefer fluconazole for treatment of cases involving the CNS, eye, or prostate because, as a consequence of its smaller molecular size and lipophilicity, it achieves better penetration into these tissues. Other reasons for choosing fluconazole include lower cost and better tolerability. Doses of at least 10 mg/kg/day are recommended.
Other azoles. Ketoconazole is infrequently used because, based on studies in blastomycosis (82), it is less effective and causes more adverse effects than itraconazole. One of five cats with histoplasmosis responded to ketoconazole (50). Posaconazole and voriconazole are more active than fluconazole and have been used successfully in humans with histoplasmosis, but have not been evaluated in animals. Histoplasma also is susceptible to isavuconazole, a newly approved antifungal azole, but clinical studies for treatment of histoplasmosis have not been reported. The newer azoles are more expensive than itraconazole or fluconazole.

**Amphotericin B.** Amphotericin B is the treatment of choice in severe cases in humans and induces a clinical response more rapidly than itraconazole (83) or fluconazole. Reasons for amphotericin B’s superiority include its fungicidal mode of action and intravenous route of administration, rapidly providing therapeutic blood concentrations. Administration of amphotericin B for the first 3 to 7 days of therapy may improve early survival, after which treatment could be changed to itraconazole or fluconazole. Lipid forms of amphotericin B are better tolerated but more expensive than the deoxycholate formulation. Renal function and serum electrolytes should be monitored during treatment.

**Adjunctive therapy.** Schulman noted rapid clinical improvement in ten dogs with mediastinal lymphadenitis causing airway obstruction, five of which also received antifungal treatment (52). Corticosteroids also may be helpful in cases of diffuse pulmonary histoplasmosis complicated by respiratory insufficiency. Concurrent antifungal therapy is recommended to reduce the risk for progressive dissemination caused by corticosteroid-induced immunosuppression.

**Monitoring Therapy.** The antigen test often is used in deciding when to stop therapy and to diagnose relapse. In 27 dogs with blastomycosis, antigen levels in urine were negative in 70% at the time of treatment discontinuation (62). (http://onlinelibrary.wiley.com/doi/10.1111/jvim.12306/abstract) Relapse occurred in 7 dogs (26%), 2 of which had a positive urine antigen at treatment discontinuation, and was associated with a rise in antigenuria in 5. The authors recommended continuing therapy until the clinical findings, including eye exam, resolved, chest radiographs were normal or stable and urinary antigen was negative. A reasonable approach would be to test for antigenuria every 3 months during therapy, at 3 and 6 months after stopping therapy, and if relapse is suspected.

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<th>Table 2. Treatment recommendations</th>
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<td>Itraconazole 5-10 mg/kg/day or fluconazole 10 mg/mg/day for 4-6 months&lt;sup&gt;1&lt;/sup&gt;</td>
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<tr>
<td>Consider amphotericin B&lt;sup&gt;2&lt;/sup&gt; in severe cases, including respiratory insufficiency&lt;sup&gt;3&lt;/sup&gt;</td>
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<td>Monitor antigen every 3 months and if suspect relapse Itraconazole blood level&lt;sup&gt;4&lt;/sup&gt;</td>
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<sup>1</sup>some cases may require more than 6 months of treatment; <sup>2</sup>lipid formulation better tolerated; <sup>3</sup>may benefit from adjunctive corticosteroid therapy; <sup>4</sup>day 14 (dogs) to 21 (cats) of therapy to ensure levels between 1.0 and 10 µg/mL and if suspect relapse or itraconazole toxicity.
Reference List


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