PROGRESSIVE CEREBELLAR ATAXIA IN AN ADULT DOG

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History
Scout, a previously healthy, 6-year-old, spayed English springer spaniel show dog, was presented with a 2-month history of progressive incoordination and head tilt. Signs progressed despite treatment with amoxicillin/clavulanic acid for presumptive otitis media or interna. Slight resolution of the head tilt was observed after an anti-inflammatory dose of prednisone. Metronidazole had not been administered.

Physical Examination
Examination showed a right head tilt, wide-based stance, nodding movement of the head and body (ie, titubation), intention tremors, ambulatory cerebellar ataxia (worse on the right), absent right menace response, mild right facial paresis, and a positional ventrolateral strabismus in the right eye. Neuroanatomical localization was to the right cerebellomedullary angle.
Scout traveled extensively before onset of signs, which started as missing an occasional step and progressed to falling after jumping, difficulty posturing to urinate, not catching objects, and head tilt.
bilateral temporal myopathy, which had not been present upon initial physical examination.

Analysis of a CSF specimen showed increased cellularity (7 nucleated cells/µL, compared with the normal 0-5 cells/µL range), increased total protein (31 mg/dL, compared with the normal <25 mg/dL range), and a mild mixed pleocytosis, possibly dampened by corticosteroid administration. The differential cell count was 43% nondegenerate neutrophils, 30% macrophages, and 27% lymphocytes (predominantly small with occasional reactive forms).

No neoplastic cells or infectious agents were detected. The results of serum and CSF indirect fluorescent antibody titers were positive for *Neospora caninum*.

**Diagnosis**

Central nervous system (CNS) neosporosis

**Treatment**

The treatment plan included administration of antiprotozoal antibiotic medications, including clindamycin (15 mg/kg PO twice a day) and trimethoprim/sulfamethoxazole (15 mg/kg PO twice a day). Although many resources recommend a 4-week course of antibiotic therapy, frequently CNS *N caninum* infections require a longer duration of treatment (eg, 6-12 months) and occasionally long-term treatment. Trimethoprim/sulfamethoxazole blocks bacterial folate synthesis, and chronic use can cause folate acid-deficient anemias; therefore, folic acid (1 mg PO once a day) supplementation was initiated.

Anti-inflammatory prednisone therapy (0.5 mg/kg PO twice a day) was tapered slowly over several months.
**Outcome**

Scout’s clinical signs deteriorated to nonambulatory tetraparesis during the first week of treatment. Two weeks after initiation of treatment, the dog began to walk and showed continued improvement during the next 7 months. Scout still hesitated to climb stairs but was able to take 10- to 20-minute walks and was otherwise healthy. The exact source of Scout’s exposure was not determined. The patient did not have exposure to cattle, but white-tailed deer and coyotes are common in the patient’s geographic area.

Neosporosis typically affects dogs younger than 6 months of age, but fatal disease can occur at any age. In puppies, 1 of the most common clinical presentations is inflammation of the muscles and nerve roots of the pelvic limbs, which causes progressive paraparesis, muscle atrophy, and loss of patellar reflexes that frequently progresses to fibrous muscular contracture with hyperextension.

In puppies and adult dogs, other organs (eg, skeletal muscles, heart, liver, skin, eyes, lungs, pancreas) may be affected. In adult dogs, various signs localized to the CNS may be seen (eg, paresis, paralysis, ataxia, head tilt, seizures). Often, these signs are multifocal. As this case illustrates, neosporosis is a differential diagnosis for progressive cerebellar ataxia in adult dogs.

![T2-weighted sagittal (A) and contrast-enhanced T1-weighted sagittal (B) MRI scans of a normal dog.](image1)

![T2-weighted sagittal (A) and contrast-enhanced T1-weighted sagittal (B) MRI scans of a 6-year-old, spayed English springer spaniel dog with CNS neosporosis. The scans depict peripheral atrophy of the cerebellum (especially dorsally) with replacement by CSF (A and B). This finding is consistent with necrotizing cerebellitis. In addition, there is contrast enhancement of the cerebellar nuclei (B). The cerebellum is circled; compare to Figure 1.](image2)
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DID YOU ANSWER?

A positive serum titer can result from infection, passive maternal transfer, or vaccination. However, canine vaccines do not exist, so a false-positive result from vaccination is unlikely. A definitive diagnosis requires identification of the parasite in CSF or tissues. A positive titer in a CSF sample is suggestive of *N caninum* (Figure 3) in the central nervous system.

The life cycle of *N caninum* involves a final host (the dog and other canids) and intermediate hosts such as mammals, birds, and cattle. In cattle, *N caninum* can cause abortions. The final canine host produces oocysts in feces that contaminate the environment. Intermediate hosts that ingest these oocysts have intracellular stages in their tissues that include tachyzoites and tissue cysts, which can infect dogs when ingested.3-6 Most canine infections that induce disease are believed to have occurred congenitally, as tachyzoites present in the dam are passed through the placenta or perhaps in the milk.

Dogs also can develop disease by eating the tissues of intermediate hosts, but the mechanism is unclear.3-6 Ingestion of sporulated oocysts by dogs rarely results in infection or further fecal transmission of oocysts into the environment. *N caninum* is not considered a zoonotic agent.

Additional Resources