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Fatal pulmonary hemorrhage associated with vascular amyloid deposition in a cat

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Abstract

Case summary An adult female spayed Siamese-cross cat of unknown age was presented for bilateral hemorrhagic otorrhea. Nasopharyngeal polyps were diagnosed by CT and biopsy; bilateral ventral bulla osteotomies were performed. Episodic epistaxis, otic hemorrhage and hemoptysis with respiratory distress progressed over 18 months. Systolic blood pressure, complete blood count, plasma biochemistries, prothrombin time, partial thromboplastin time and coagulation factor 12, 9 and 8 activities were normal. Serial thoracic radiographs revealed patchy interstitial to alveolar patterns. Airway hemorrhage prevented diagnostic bronchoscopy. Respiratory hemorrhage was ultimately fatal. Amyloid deposition was identified in pulmonary vasculature, bronchial wall, lymphoid tissues, nasal-pharyngeal tissue and tympanic bullae based on microscopic examination and confirmed by Congo red staining with green birefringence under polarized light.

Relevance and novel information Amyloidosis should be considered as a differential diagnosis in cats with spontaneous hemorrhage of the respiratory or otic tracts. Although systemic amyloidosis is associated with a grave prognosis, this case suggests that prolonged survival is possible after the initial onset of signs in cats with pulmonary amyloidosis.

Keywords: Serum amyloid A; hemoptysis; otorrhea; epistaxis; nasopharyngeal polyps; otic hemorrhage

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Case description

An adult female spayed indoor-only Siamese-cross cat of unknown age was presented for hemorrhagic otorrhea (time 0). The cat had been adopted from an animal shelter as an adult 4 years prior; at which time complete blood count, biochemistry panel, urinalysis and feline leukemia virus/feline immunodeficiency virus testing were unremarkable. Four months after adoption, the cat developed yeast otitis externa of the left ear. Although otitis resolved with treatment (topical clotrimazole/dexamethasone solution q12h for 14 days), head shaking progressed and was accompanied by mild serous nasal and ocular discharge. ACT scan of the ears and video otoscopy were performed, which were unremarkable, and untargeted nasal biopsies revealed mild lymphocytic and histiocytic rhinitis. Clinical signs resolved without treatment.

Diagnostics performed and therapeutics implemented after re-presentation (time 0) are summarized in Table 1. Briefly, evaluation revealed spontaneous bilateral hemorrhagic otorrhea, fluid-filled bullae, bilateral ruptured tympanic membranes and a mass in the left aural canal. The mass was removed using manual traction and diagnosed as a nasopharyngeal polyp based

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Table 1 Chronological summary of clinical signs, physical examination abnormalities, diagnostic findings and treatments in a cat with progressive spontaneous episodic hemorrhage of the respiratory and otic tracts

Time point	Presenting complaints and abnormalities on examination	Diagnostics performed	Abnormalities on diagnostic evaluation	Treatment
0	Bilateral hemorrhagic otorrhea	CBC*, plasma biochemistry panel†, PT‡, PTT‡, CT scan Mass removal by manual traction Nasal biopsies with histopathology	Ruptured tympanic membranes Nasopharyngeal polyp in the left ear	Prednisone (5 mg PO q24h for 7 days then q48h for 7 days) Topical flucinolone with dimethyl sulfoxide and topical silver sulfadiazine for both ears
1 month	Bilateral hemorrhagic otorrhea	Histopathology	Bilateral nasopharyngeal polyps	Bilateral ventral bulla osteotomies
6 months	Intermittent epistaxis Gingival recession Stage 1 periodontal disease	BP§ 170–180 mmHg (with acclimation decreased to 140–150 mmHg) Ophthalmic/fundic examination CBC, plasma biochemistry panel PT, PTT Dental radiographs Otic and oral exams under general anesthesia	Blood noted on endotracheal tube upon extubation	Dental cleaning
8 months	Intermittent hemoptysis and epistaxis	BP 150 mmHg Thoracic radiographs Coagulation factor 8, 9 and 12 analysis	Interstitial infiltrate in caudoventral aspect of left cranial lung lobe Ill-defined nodule in right caudal lung lobe	None
8.5 months	Intermittent hemoptysis and epistaxis	Thoracic radiographs CBC, plasma biochemistry panel Blastomyces urine antigen quantitation	Interstitial infiltrate unchanged in left cranial lung lobe Nodule in right caudal lung lobe more defined and another soft tissue opacity in same lung lobe	Itraconazole (10 mg/kg PO q24h) for 2 weeks
9.5 months	Intermittent hemoptysis and epistaxis Grand mal seizure	Thoracic radiographs MRI of head CSF tap for cytology Video otoscopy Bronchoscopy	Mild interstitial pattern in right caudal lung lobe Interstitial pattern in caudal subsegment of left cranial lobe Right tympanic bulla and ear canal filled with suspected hemorrhage Bulging tympanic membrane in the right ear Tracheal hemorrhage	Myringotomy with removal of blood clot in the right ear Tobramycin otic drops for both ears
10–18 months	Intermittent hemoptysis with respiratory distress			Prednisolone (5 mg PO q24h) Enrofloxacin (5 mg/kg PO q24h for 10 days) Phenoxybenzamine (2.5 mg PO q12h)

(continued)

Table 1 (Continued)

Time point	Presenting complaints and abnormalities on examination	Diagnostics performed	Abnormalities on diagnostic evaluation	Treatment
18 months	Intermittent hemoptysis with respiratory distress	BP 130 mmHg Sedation (butorphanol [0.4 mg/kg] and dexmedetomidine [2 µg/kg IV]) Thoracic radiographs Limited abdominal ultrasound CBC, plasma biochemistry panel <i>Toxoplasma</i> IgG and IgM antibody titers	Multifocal, coalescing, structured, interstitial and alveolar lung patterns. Pleural effusion Small adrenal glands Non-regenerative normocytic normochromic anemia (hematocrit 34%) <i>Toxoplasma</i> -positive IgG titer (≥ 8192) and negative IgM titer (< 512)	
18.3 months	Acute hemoptysis and severe respiratory distress	Necropsy	Severe chronic-active widespread multifocal lymphoplasmacytic, suppurative and hemorrhagic broncho-interstitial pneumonia with bronchiectasis, and extensive fibrosis. No infectious organisms seen on routine or gram special staining Amyloidosis type AA involving the pulmonary vessels, bronchi, bronchioles, retropharyngeal lymph node and tympanic bullae	Terbutaline (0.01 mg/kg SC) Butorphanol (0.2 mg/kg SC)

*ADVIA 2120i (Siemens Medical Solutions USA)

†Cobas c 501 analyzer (Hoffman-LaRoche)

‡STA Compact, Diagnostica Stago

§Indirect systolic blood pressure (BP) taken using the Doppler flow method

CBC = complete blood count; PT = prothrombin time; PTT = partial thromboplastin time; CSF = cerebrospinal fluid

on anatomic location, gross appearance and histologic changes. Intermittent bilateral hemorrhagic otorrhea persisted over the next month in spite of medical therapy. Sedation and/or analgesia was administered as appropriate for all described diagnostic and therapeutic procedures. Bilateral ventral bulla osteotomies were performed to remove bilateral nasopharyngeal polyps. Episodic hemorrhagic otorrhea continued.

Five months postoperatively, unilateral epistaxis of unknown cause occurred during physical examination for dental prophylaxis. An etiology was not identified on diagnostic testing, although primary hemostasis was not evaluated. Dental prophylaxis was uneventful, but blood was noted on the endotracheal tube upon extubation.

Hemoptysis developed 2 months later, which increased in frequency over the next 6 weeks. Diagnostic findings included multifocal shifting infiltrates and nodular change on radiographs (Figure 1).

Neurologic evaluation, including MRI and cerebrospinal fluid analysis, was performed owing to a grand mal seizure 6 weeks after the onset of hemoptysis. The right tympanic bulla and ear canal were filled with heterogeneously hyperintense material on T2-weighted sequences consistent with hemorrhage. A large blood clot was removed via myringotomy. Bronchoscopy was attempted, but extensive hemorrhage in the trachea precluded visualization. Given the patient's instability and our inability to visualize the tracheal lumen due to hemorrhage, bronchoalveolar lavage was not attempted. Recovery from anesthesia was complicated by hypoxemia and repeated obstruction of the endotracheal tube with blood. The cat was discharged the following day.

Over the next 9 months, hemoptysis recurred with increasing frequency and severity, although the cat was clinically normal between episodes. Empiric treatment trials had no apparent effect on clinical signs (Table 1).

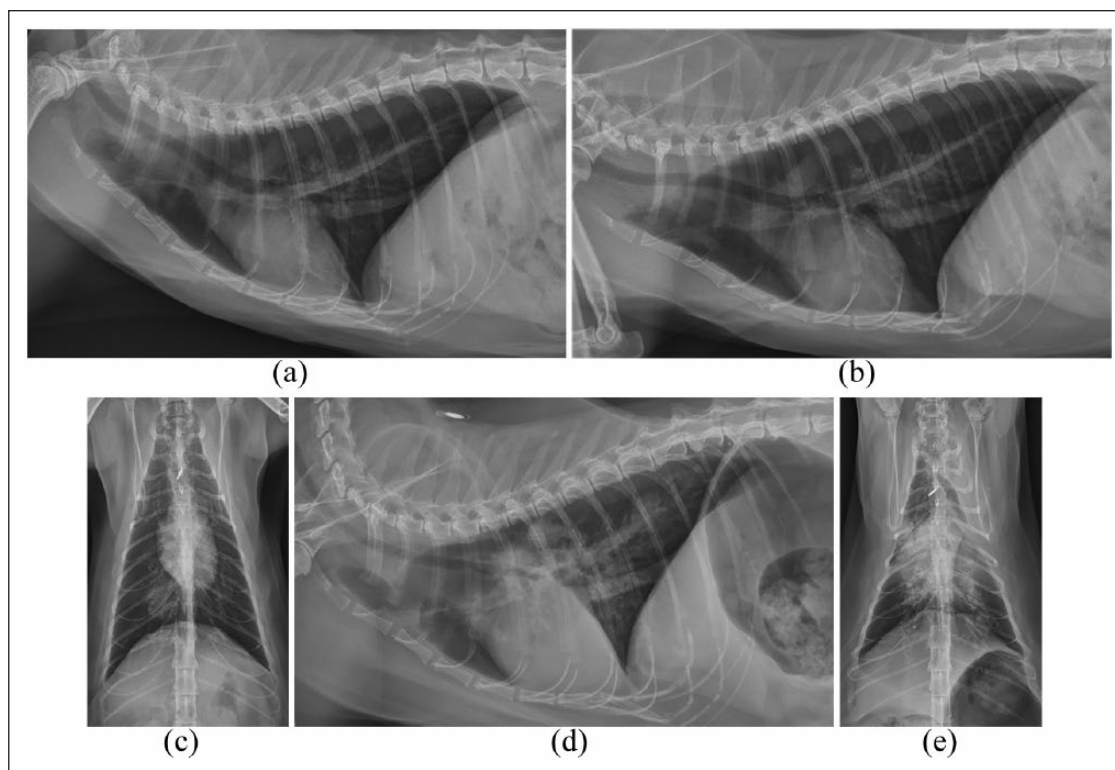


Figure 1 Multifocal shifting infiltrates and nodular change on thoracic radiographs taken at 8.5 months ([a] right lateral view, [b] left lateral view, [c] ventral/dorsal view) and 18 months ([d] right lateral view, [e] ventral/dorsal view) in a cat with episodic spontaneous hemoptysis.

Re-evaluation at the hospital resulted in epistaxis, respiratory distress and temporary oxygen dependency. Ten days later, the cat was presented for acute hemoptysis with severe respiratory distress and died 5 h after placement in an oxygen cage.

At necropsy, there was hemorrhagic discharge from the nares and free blood within the nasal passages. Lung lobes had multifocal-to-extensive reddening with palpable firmness, particularly in the hilar region. The bronchi were filled with yellow-to-green mucoid material. Unfortunately, only grossly abnormal organs (respiratory system, retropharyngeal lymph nodes, liver and adrenal glands) were evaluated microscopically.

On histologic examination, bronchioles were variably ectatic and filled with epithelial and neutrophilic cellular debris, red blood cells and mucus. This inflammatory infiltrate extended into the immediately adjacent alveolar lumens and septae. In addition, hemorrhage comprised 25–95% of the lung sections, blending with the inflammation and occasionally obliterating the surrounding parenchyma (Figure 2). The walls of pulmonary vessels, bronchi and bronchioles in these areas were segmentally variably distorted and expanded by scant-to-abundant amorphous, eosinophilic, hyaline material. This material was congophilic, potassium permanganate sensitive, displayed apple green birefringence when polarized and positive for anti-AA antibody reactivity on

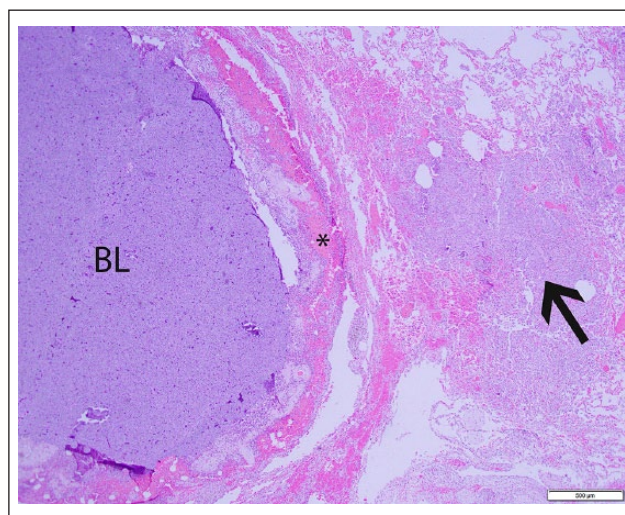


Figure 2 Debris-filled bronchiectatic airway lumen (BL) with bronchial wall disruption, peribronchial-to-alveolar hemorrhage (*) and adjacent localized pneumonia (black arrow). Hematoxylin and eosin stain. Bar = 500 μm

immunohistochemistry performed using Marmoset AA 408-89 monoclonal antibody (Immunology and Cancer Program, University of Tennessee Graduate School of Medicine, Knoxville, TN, USA) as per previously described techniques (Figures 3 and 4).¹ Submucosal and

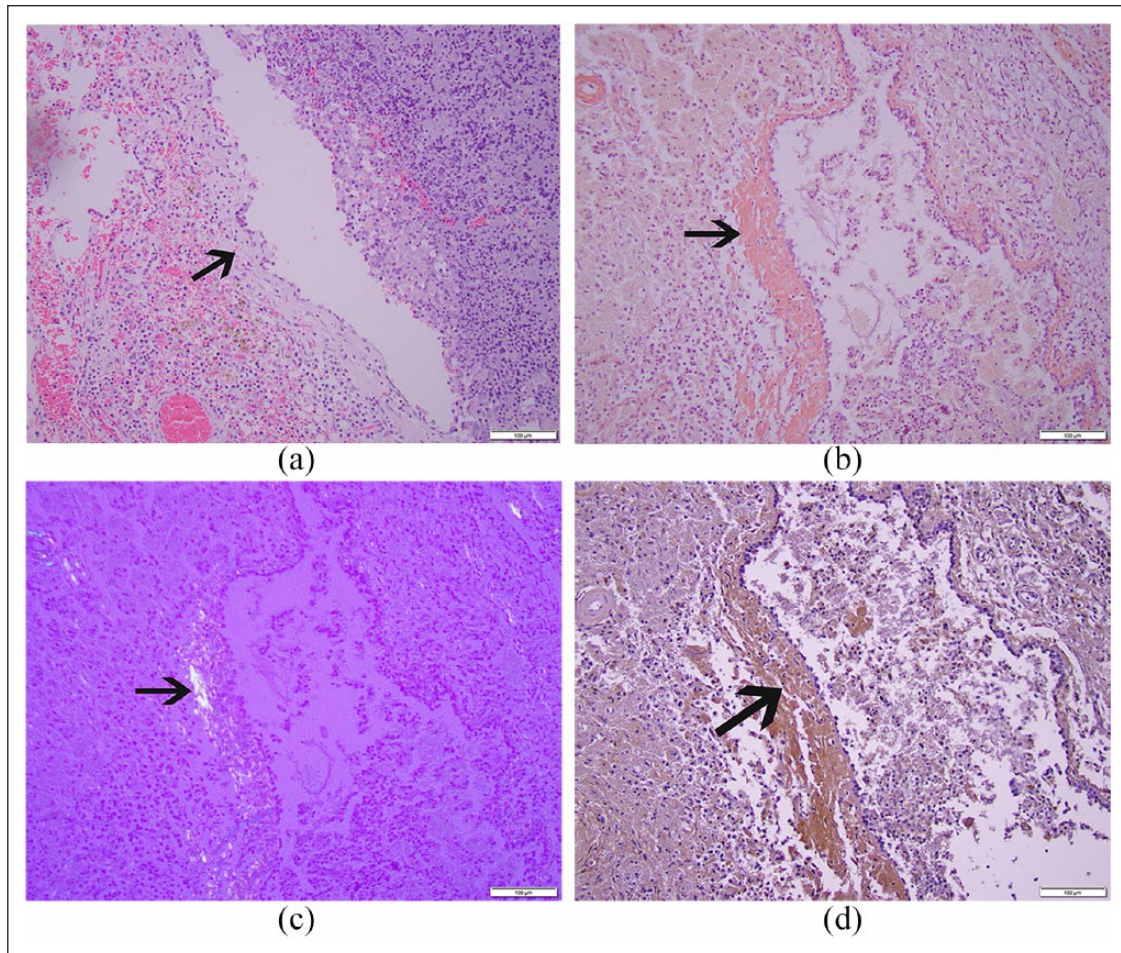


Figure 3 Bronchial wall tissue section with mural amyloid deposition (arrow): (a) hematoxylin and eosin stain, (b) Congo red stain without and (c) with polarization, and (d) positive immunohistochemical staining for AA antibody. Bar = 100 µm

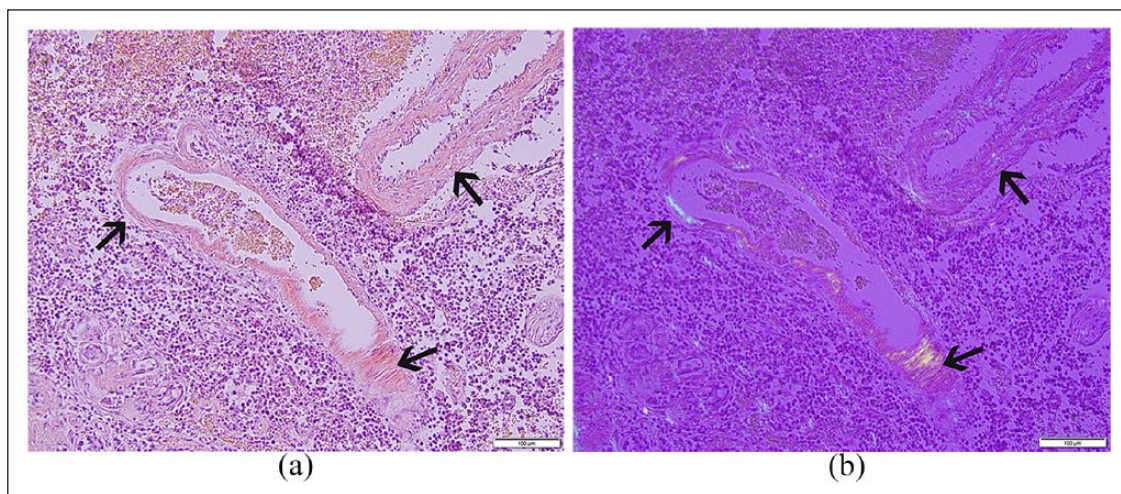


Figure 4 Pulmonary vessels with mural amyloid deposition (arrows). Congo red stain (a) without and (b) with polarization. Bar = 100 µm

arteriolar amyloid also was identified in the retropharyngeal lymph node and archived biopsies from the nasopharyngeal polyps and tympanic bullae.

On gross examination, the liver was small (1.78% of body weight vs 3–3.5% normal). On hematoxylin and eosin-stained sections, the tunica media of large hepatic

arteries was segmentally mildly expanded by smudgy pale eosinophilic material. However, periarteriolar birefringence, not anomalous color birefringence, was identified on special stains of these tissues. Thus, this infiltration was consistent with collagen deposition. Final diagnoses for the cat were pulmonary amyloidosis type AA, bronchiectasis and pneumonia with fibrosis and extensive pulmonary hemorrhage.

Discussion

This report describes a novel presentation of amyloidosis type AA in a cat. Amyloidosis is a non-specific term referring to a group of disorders caused by extracellular *in vivo* deposition of insoluble fibrillar proteins, such as serum A-amyloid protein (type AA), in a beta-sheet conformation.^{2,3} Many types of amyloidosis originating from different proteins have been described in people – some of which are inherited. Amyloid deposition in the extracellular space disrupts the normal tissue architecture. Although organ function can be significantly affected by the site and distribution of amyloid deposition, the mechanisms are incompletely understood.^{4,5} Simple physical, mechanical replacement of parenchyma by amyloid deposits might not be sufficient to cause dysfunction. Once formed, the amyloid is relatively insoluble and is resistant to proteolysis. Treatment is aimed at stopping the underlying disease process if possible.

Amyloidosis in the cat most commonly involves the kidney and pancreatic islets.^{6,7} Familial amyloidosis (type AA) has been recognized in Abyssinian cats, resulting primarily in renal deposition and disease.^{7,8} No biochemical, urinalysis or gross necropsy findings consistent with renal disease or diabetes mellitus were identified in this case. Unfortunately, owing to their normal gross appearance, renal and pancreatic tissues were not retained for histologic examination. As such, subclinical renal and pancreatic amyloidosis cannot be ruled out. Siamese and other breeds can develop type AA systemic amyloidosis, primarily depositing in the sinusoids of the liver with potential resultant spontaneous liver rupture.^{9–13} Most cases with hepatic amyloidosis and rupture have increased serum alanine aminotransferase activity, anemia and grossly abnormal livers on necropsy.^{10,13} Although the cat of this report was a Siamese-cross, plasma liver enzyme activities and coagulation tests were normal, the liver was grossly small in size and there was no evidence of hepatic sinusoidal amyloid deposition on special staining. Because amyloid sequencing was not performed, a genetic component cannot be ruled out.

Clinically significant pulmonary amyloidosis has not been previously reported in the cat. In 4/11 Abyssinian cats with renal amyloidosis in one report, amyloidosis was found in limited areas of the peribronchial and peribronchiolar connective tissue.¹⁴ Pulmonary amyloid was not identified in other feline reports of amyloidosis.^{9–12,14} Amyloidosis has been associated with bronchiectasis in

humans, but amyloid deposition and disease is primarily renal in those cases.¹⁵ In comparison, examination of histologic samples from the lungs of six cats diagnosed with bronchiectasis at the time of necropsy did not identify amyloid in the bronchiolar or pulmonary vessel walls (RL Donnell, unpublished data).

The cat of this report had histopathological evidence of bronchopneumonia and bronchiectasis, possibly secondary to recurrent airway hemorrhage and/or disruption of the mucociliary apparatus due to amyloid deposition. The lack of clinical signs between hemorrhagic episodes and the slow clinical progressive nature of the condition could reflect chronic inflammation leading to bronchiectasis and amyloidosis. Alternatively, the deposition of amyloid in bronchial walls and pulmonary vasculature caused the inflammation, bronchiectasis and hemorrhage. In humans, severe-to-fatal hemorrhage due to amyloidosis has been described with amyloid deposits in intestines, urinary bladder, mediastinum, trachea, bronchi and larynx, as well as cerebral arterioles.^{16–21} Individuals with tracheobronchial, laryngeal and mediastinal amyloidosis typically present with respiratory signs due to mechanical obstruction from localized amyloidosis and resultant hemorrhage.

Vascular mural amyloid deposition with pulmonary hemorrhage has been reported in people.^{16,22–24} Idiopathic pulmonary vascular amyloidosis, occasionally resulting in respiratory distress due to diffusion impairment, has been described in older dogs secondary to apolipoprotein AI deposition.^{25–27} In contrast, clinical signs in this cat were due to acute recurrent hemorrhage with apparently normal pulmonary function between episodes. Alterations in vascular compliance due to the severity of amyloid type AA deposition in the walls, potentially exacerbated by increased pulmonary artery pressures during times of stress, likely resulted in intermittent pulmonary hemorrhage. Chronic topical steroid exposure is reported to contribute to the occurrence of telangiectasia in humans and experimental rats.^{28,29} As such, it is possible that administration of steroids contributed to the final fatal hemorrhagic event.

Conclusions

This case report describes a novel presentation of systemic amyloidosis type AA in a cat. Amyloid deposition was primarily concentrated in the walls of the pulmonary vasculature and bronchioles with lesser involvement of the nasopharyngeal tissues and local lymph nodes. Clinical signs were limited to spontaneous, progressive hemorrhage of the otic and airway tracts, and hemorrhage was ultimately fatal. Final diagnosis of amyloidosis was achieved after using microscopic examination and Congo red staining with green birefringence under polarized light on necropsy tissues. Amyloid deposition was subsequently confirmed in archived biopsy specimens from early in the disease course.

Based on the findings in this case, systemic amyloidosis should be considered as a differential diagnosis in cats with spontaneous hemorrhage of the respiratory or otic tracts after other causes have been eliminated. Evaluation of special stains on biopsy specimens is required to achieve a diagnosis during the early phase of the disease. This case shows that, although long-term prognosis for systemic amyloidosis is grave, good quality of life is possible in some cats with pulmonary amyloidosis due to the slowly progressive nature of clinical signs.

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