Hepatic fibrosarcoma incarcerated in a peritoneopericardial diaphragmatic hernia in a cat

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Abstract

Case summary A 14-year-old, female neutered domestic shorthair presented for dyspnoea. Thoracic ultrasonography and radiography showed that a heterogeneous mass was present within the pericardial sac, and the mass continued caudally with the mesenteric fat. On CT, the outline of the diaphragm was not continuous and there was an obvious defect with diaphragmatic thickening present at the mid-level of the liver. A pleural effusion and a small-volume pericardial effusion were also present. A ventral midline celiotomy and median sternotomy revealed a 5 × 6 × 7 cm firm, irregular, tan-coloured soft tissue mass within the pericardial sac attached to both the diaphragmatic defect and liver. The mass was carefully dissected away from the heart and the diaphragmatic defect was repaired with primary closure. Postoperatively, the cat had a persistent pneumothorax that required continuous pleural suction for 41 h. The cat died 44 h postoperatively. Histopathology and immunohistochemistry confirmed the mass to be a hepatic fibrosarcoma incarcerated in a peritoneopericardial diaphragmatic hernia (PPDH).

Relevance and novel information This is the first reported case of metaplastic transformation of liver into a sarcoma in a cat with PPDH. In addition, hepatic fibrosarcoma is a rarely reported location for fibrosarcoma in this species.

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Introduction

Fibrosarcoma is a common cutaneous neoplasm of cats, representing 15% of cases in one study of 340 cats.1 The hepatic form is reported much less frequently.2,3 One series documented hepatic fibrosarcoma in only 1/41 cases of non-lymphomatous hepatobiliary masses.3 Peritoneopericardial diaphragmatic hernia (PPDH) is a structural abnormality resulting in communication between the pericardial sac and peritoneal cavity. The embryogenesis of PPDH is unknown but may include malformation of the pleuroperitoneal folds and septum transversum.4–6 It is often discovered as an incidental finding in cats and, consequently, when clinical signs are absent, surgical correction is difficult to justify.7,8

Previous case reports have documented a possible association between incarcerated hepatic tissue in PPDH and the development of hepatic myelolipomas and hepatic cysts.6,9–12 This report describes a hepatic fibrosarcoma incarcerated in a PPDH in a cat. To our knowledge no such association has previously been documented in this species.

Case description

A 14-year-old, female neutered domestic shorthair (body weight 4.5 kg) was referred following a 1 day history of dyspnoea. The cat spent time outdoors, but no history of trauma was reported. The cat had a stable body weight. Immediately prior to referral a cumulative dose of furosemide 8 mg/kg was administered intramuscularly.

On physical examination the cat was hypothermic (36.7°C), had an inspiratory dyspnoea, reduced lung...

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sounds, tachypnoea (respiratory rate 52 breaths per minute), a non-compliant thorax and was tachycardic (heart rate 240 beats per minute); left parasternal gallop sounds were auscultated. Oxygen saturation was 88%. Systolic blood pressure measured from the left forelimb was 150 mmHg using a size 3 cuff (Doppler; Parks Medical Electronics).

Inspiratory dyspnoea without upper airway noise is most commonly due to pathology of the pleural space (effusions, pneumothorax). Owing to the cat’s age and presenting clinical signs, pleural effusion due to cardiac failure, with or without primary causes (hyperthyroidism, hypertension) or neoplasia were suspected most likely. The degree of respiratory compromise warranted that diagnostic tests to differentiate these differentials were performed first. These included dorsoventral radiography of the thorax and a brief transthoracic ultrasound (CX50; Philips) while the cat was in sternal recumbency. Mask inhalation oxygen was administered during these procedures. No sedation was required.

A bilateral pleural effusion was noted radiographically and was anechoic in appearance ultrasonographically. For diagnostic and therapeutic reasons, ultrasound-guided thoracocentesis was performed and 250 ml of serous fluid was removed. Protein concentration of the effusion was 37 g/l (reference interval [RI] 0–25 g/l). Nucleated cell count was 1.7 × 10^9/l (RI 0–1.5 × 10^9/l). Red cell count was 0.1 × 10^12/l (RI 0 × 10^12/l). These values are consistent with a modified transudate. Cytology and cell counts of the fluid revealed 24% mature neutrophils, 14% small lymphocytes and 62% vacuolated and phagocytic macrophages. No other significant cell population or microorganisms were identified.

Echocardiography (ECG; right parasternal) demonstrated a 1 mm pericardial effusion. Hypertrophy of the myocardium (left ventricular wall at end diastole 7.5 mm [normal <5.5 mm]) was evident. No other ECG abnormalities were noted. Specifically, the left atrium: aorta diameter ratio was 1.1 (RI <1.5). There was no ultrasonographic evidence of cardiac tamponade. Transthoracic ultrasonography identified a 4.27 cm heterogeneous mass (measured in a medial to lateral plane) caudal to the heart (Figure 1). The mass appeared to be closely associated with the diaphragm. Abdominal ultrasound revealed diminished hepatic volume. Abdominal hepatic tissue was homogeneous.

A mild non- or pre-regenerative anaemia (haematocrit 26% [RI 30.3–52.3%]) was present on haematology screening (IDEXX ProCyte Dx Haematology Analyser). A stress leukogram was present. Red and white cell morphology was normal.

Serum biochemistry (IDEXX Catalyst Dx Chemistry Analyser) revealed a mild hypokalaemia 3.6 mmol/l (RI 3.7–5.4 mmol/l), mild hyperglycaemia 11.0 mmol/l (RI 3.2–7.6 mmol/l), mild azotaemia (urea 18.0 mmol/l [RI 5.0–15.0 mmol/l]) and a mild panhypoproteinaemia (albumin 24 g/l [RI 25–38 g/l] and globulin 30 g/l [RI 31–52 g/l]). Total thyroxine (T4) was within the normal range (44 nmol/l [RI 10–60 nmol/l]). Urine specific gravity (USG) was isostenuric (1.010).

The patient’s azotaemia, hypokalaemia and isostenuria was considered to be caused by a combination of reduced appetite and thirst prior to presentation, followed by dehydration from furosemide administration. Mild chronic renal failure was another differential; however, the recent diuretic administration made the interpretation of USG as ‘appropriate’ or not difficult to determine.

CT was performed (Aquilion 64; Toshiba Medical Systems) after the cat was sedated with methadone 0.9 mg SC (Methone; Ceva), induced with alfaxalone 5 mg IV (Alfaxan; Jurox Pty) and oxygen supplementation provided. Iohexol (Omnipaque; GE Healthcare) 1800 mg IV was administered for the postcontrast series. The CT scan demonstrated that the intrathoracic mass was continuous with the intra-abdominal lobes of the liver and extended cranially through a large diaphragmatic defect into the pericardial space (Figure 2). The mass (7 cm) had a heterogeneous pattern of enhancement compared with the homogeneous intra-abdominal hepatic tissue. Lymphadenopathy of the sternal lymph nodes was evident.

A diagnosis of a PPDH was made. Differentials for the intrathoracic mass included granulomatous disease of viral and non-viral causes, migrating foreign body, bacterial empyema with abscessation and neoplasia (of hepatic and non-hepatic origin). The sternal lymphadenopathy was presumed to represent localised inflammation or metastasis.
A cranioventral midline celiotomy was performed in order to repair the diaphragm and remove the intrathoracic lesion, if possible. The cat was premedicated with methadone 0.9 mg SC and induced with alfaxalone 5 mg IV. Fentanyl 10 μg/kg/h IV constant rate infusion (CRI) (DBL Fentanyl Injection; Hospira) was administered throughout the surgery. Isoflurane (0.5–1.5%) was used for maintenance. Lactated Ringer’s solution (Hartmanns Compound Sodium Lactate; Baxter Viaflex) was administered at a rate of 5–10 ml/kg/h.

The coeliotomy was extended cranially and a median sternotomy was performed using a sagittal saw. A 3–4 cm radial diaphragmatic hernia was identified at the xiphoid cartilage of the sternebrae, within which the liver lobe (right lateral) and herniated omentum were present. The hernia contents were manually reduced into the abdomen. A biopsy of the left liver lobe (present in the abdomen) was taken using the guillotine method with 3/0 Polydiaxonone (PDS; Ethicon Johnson & Johnson).

A 5 × 6 × 7 cm firm, irregular, tan-coloured soft tissue mass was visualised surrounding the heart on the right side and within the pericardial cavity (Figure 3). There were satellite nodules of similar appearance adherent to the pericardium (considered possible metastases). These were biopsied. The mass was adhered to the diaphragmatic defect and liver by fibrous strands. The heart was free of macroscopic tumour. The mass was carefully resected with blunt and sharp dissection and bipolar cautery was used to maintain haemostasis. A subtotal pericardectomy (subphrenic) was performed to reduce the risk of a pericardial effusion redeveloping at a later period.

The diaphragmatic hernia edges were debrided and closed with continuous 2/0 polydiaxonone. The medial sternotomy was closed with a single 0.8 mm orthopaedic wire (Roth Medical) and multiple 1/0 polydiaxonone loops around the sternebrae. Polydiaxonone was used owing to insufficient orthopaedic wire available at the time of the procedure. The remaining thoracotomy and laparotomy incision was closed in a three-layered approach. Specifically, the linea alba and subcutaneous tissue was sutured with 2/0 and 3/0 PDS in a simple continuous pattern and skin with 3/0 nylon (Riverlon; Riverpoint) in a forward interlocking pattern. Two drains (SurgiVet Chest Drainage Tubes, 12 Fr 42 cm) were placed in the dorsolateral aspect of the eighth intercostal space at the left and right thoracic wall. Negative thoracic pressure was achieved. A 14 Fr oesophageal feeding tube was placed (Esophagostomy Tube, Feline [14 Fr, 33 cm]; MILA International).

The cat required mechanical ventilation throughout the anaesthesia. Intraoperative ECG revealed one short period of ventricular tachycardia (30 s) and the occasional ventricular premature complex that did not warrant therapy. Anaesthesia was otherwise uneventful. Postoperative analgesia was provided with a combination of a fentanyl intravenous CRI (5–7.5 μg/kg/h) and...
bupivacaine 5 mg (Marcain; AstraZeneca) diluted with 0.9% sodium chloride via the chest drain q8h for analgesia. For a short period (15 mins) a higher dose of fentanyl (20 μg/kg/h) was used to limit excitement on recovery. Cephazolin 100 mg IV q8h (Cephazolin Sandoz; Sandoz) was administered for antibiotic coverage. Lactated Ringer’s solution, supplemented with 30 mmol potassium chloride per litre, was infused intravenously at 20–30 ml/h.

Two hours postoperatively the cat became dyspnoeic with open-mouth breathing. Oxygen saturation was between 85% and 90%. Thoracic radiography revealed bilateral pneumothorax. Subsequently, intranasal oxygen was administered and continuous suction drainage (Thora-seal iii; Medline) from the pleura performed for approximately 41 h. At this time, the pneumothorax appeared to resolve and oxygen saturation remained >95% without pleural suction. Over 41 h, 75 ml serosanguinous fluid was removed from the pleural space. A continuous ECG identified the occasional ventricular premature complex that did not warrant therapy. Forty-four hours postsurgery the cat was found to be deceased; 5 mins prior to death the cat was noted to have normal vital signs and an oxygen saturation of 96%.

The owners declined post-mortem examination. We speculate that either recurrence of pneumothorax or a cerebrovascular accident were the most likely differentials for the cause of death.

Histological examination of the mass revealed a malignant, markedly pleomorphic spindle cell proliferation that invaded and compressed small regions of hepatic parenchyma embedded within the mass (Figure 4). Some of the cells formed a collagenous matrix (confirmed with Van Gieson’s staining) (Figure 5); however, the majority showed no differentiation. Cytoplasm was a moderate-to-dark eosinophilic grey colour. Nuclei exhibited marked anisokaryosis, with bizarre round-, ovoid-, triangular-, cigar- and spindle-shaped nuclei observed. The chromatin ranged from dark smudged to pale and vesiculate, and some had prominent and multiple nucleoli. The mitotic rate was 12 per 10 high-power fields (HPFs) and there were many bizarre mitoses.

Isolated islands of hepatic tissue within the mass had essentially normal hepatic architecture, although there was invasion of the sinusoids by the malignancy (Figure 6).

Sections of adjacent pericardium were also examined, revealing moderate reactive pericarditis. Sections of non-herniated normal liver showed a moderate centrilobular fatty change and mild chronic cholangitis, with no evidence of a neoplasm.

Immunohistochemical staining was performed to determine the origin of the tumour using antivimentin antibody (stains mesenchymal, mesothelial and round cells [hepatocytes stain negative]) (Figure 7). Antipan-cytokeratin antibodies (AE1/AE3), which stain epithelial cells and well-differentiated bile ducts, were used
instead of CK19 as a feline-validated CK19 marker could not be sourced in Australia. One hundred percent of the proliferating spindle cells stained strongly positive for vimentin. The proliferating cells in the mass were negative for AE1/AE3, although multifocal sparsely distributed bile ducts within the mass stained strongly positive (Figure 8). The differentials mesothelioma or scirrhous hepatocellular/biliary carcinomas could be excluded because the proliferating cells were AE1/AE3-negative. In the mass the proliferative marker Ki-67 was positive in 12 per every 100 cells examined.

In comparison, staining of sections of pericardium (location of satellite nodules) revealed that 50% of non-neoplastic pericardial mesothelial cells were positive for AE1/AE3. This is an expected staining pattern for mesothelium.

A combination of histological morphology of the neoplasm and immunohistochemical properties demonstrated that the vast majority of proliferating cells were mesenchymal in origin, warranting the diagnosis of sarcoma. As there was a significant component of collagen production (confirmed with Van Gieson’s staining) the mass was classified as a fibrosarcoma.

Our clinicopathological assessment was that the cat had a PPDH with associated liver lobe incarceration. This liver lobe subsequently underwent neoplastic transformation to produce a hepatic fibrosarcoma.

Discussion
Feline fibrosarcoma is a malignant neoplasia that in some cats is associated with sites of inflammation, particularly vaccination injection sites. Cats have an increased tendency to respond to inflammation by fibroblast proliferation and, ultimately, tumour genesis compared with other species. Transition zones that pass from inflammation to sarcoma have been microscopically observed and support this theory. Hence, it is thought that any source of inflammation can lead to sarcoma formation in the cat. Occasionally, the tumour appears spontaneously without a known source of inflammation. Hepatic fibrosarcoma is one such example (albeit rare) in the cat.

In cats intrapericardial hepatic cysts and myelipomas have been linked with PPHA in numerous case reports. In these reports it has been speculated that incarceration of hepatic tissue in the PPDH led to vascular and lymphatic congestion and hypoxia of hepatic tissue predisposing to formation of cysts or myelipomas. In people, it has been suggested that as a result of incomplete vascular and/or ductal systems, ectopic livers are predisposed to hepatocarcinogenesis due to longer exposure of ectopic liver tissues to carcinogenic substances. The authors of this mentioned study speculate that incarceration of liver led to vascular and lymphatic constriction predisposing to neoplastic transformation to hepatic fibrosarcoma. Presumably, the mechanisms described above would have the potential to elicit neoplastic transformation in hepatic mesenchymal cells, as well as hepatocytes.

In this case, multiple negative prognostic markers were apparent on histopathology and immunohistochemistry. Specifically, nuclei exhibited marked anisokaryosis, with multiple bizarre nuclei identified within cells. The mitotic rate (12 per 10 HPFs) was high. In canine sarcomas, mitotic rate index was prognostic for development of metastasis. Additionally, in this case, the high Ki-67 index of the mass could have been associated with a poor long-term prognosis.
cats that had signs of lymphatic invasion and those that survived <1 year after surgery. In retrospect, the enlarged sternal lymph node could have been aspirated or excised intraoperatively, to assess for malignancy.

We believe the myocardial hypertrophy identified on ECG was an incidental finding given there was no evidence of heart failure on ECG or radiography. Dehydration was considered the most likely cause. Hyperthyroidism could not be totally excluded as a cause of myocardial hypertrophy given the T4 was at the upper end of the RI, in the presence of euthyroid illness. Primary hypertrophic cardiomyopathy was considered the other likely differential for the myocardial hypertrophy.

The exact cause of death in this cat will remain unknown as post-mortem examination was declined. Complications that result in severe clinical deterioration and mortality following intrathoracic surgery include pneumothorax, re-expansion pulmonary oedema, haemorrhage and malignant cardiac arrhythmias. Additionally, cerebrovascular accident is a reported postoperative complication in people and cats and therefore should be considered as a plausible cause of patient mortality. Although these complications do not necessarily result in death in every instance, animals undergoing thoracic surgery may be further compromised by changes in breathing patterns brought on by surgery, narcotic pain relief and recumbency, thus affecting their ability to handle these challenges. In our opinion, the per-acute nature of the cat’s deterioration makes pneumothorax or cerebrovascular accident the most likely possibilities.

Specifically, pneumothorax has been reported following diaphragmatic hernia repair in dogs and cats. Apparent or non-apparent iatrogenic lung lacerations may seal temporarily only for the laceration to open spontaneously during recovery as respiratory excursions and patient mobility increase. In this cat, although it appeared that the pneumothorax had resolved, premature discontinuation of active pleural suction may have lead to a fatal recurrent pneumothorax.

Although single isolated periods of ventricular tachycardia were identified intra- and postoperatively, ECG recording immediately prior to the cat’s death demonstrated normal sinus rhythm, suggesting that a malignant arrhythmia was an unlikely cause of death. Exsanguination and pulmonary oedema were improbable as intraoperative haemorrhage was not appreciated during surgery, nor were there any markers of haemorrhage or pulmonary oedema on examination shortly before death.

**Conclusions**

This is the first reported feline case of PPDH where metaplastic transformation of hepatic tissue into a fibrosarcoma has occurred. While uncommon, this should be included in the list of differentials in cats with chronic diaphragmatic hernia presenting with pleural effusions and/or thoracic mass lesions.

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