



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

PRESENTING CLINICAL SIGNS

Hx from 7/14: - Initially presented w/anorexia, ADR, v/d for past few weeks - no improvement - BMs softer than normal - 2 days ago, meds finished - not interested in food, sleeping a lot, wobbly on feet; vomited white frothy liquid 3-4x Previous/Current Meds/Last Dose: 1) Cerenia/nausea/7-10-2022 2) Metronidazole/antibiotic/7-10-2022 3)Clavamox/antibiotic/7-11-2022 (morning) 4) Gabapentin/pain control/within last 3 days (7-12 to 7-14) 5) Restoralax/stool softener/7-15-2022 (ceased, due to stool softening)

Abnormal PE/Chem/CBC/UA Results: 07-03-2022 (results listed below in following order:)
Test/Result/Flag/Normal/Range/Measure ProCyte_Dx 07-03-22 11:17p EOS 0.12 0.17 1.57 x10⁹/LL
Catalyst One 07-03-22 11:34p CREA 61 71 212 µmol/LL GLOB 54 28 51 g/LH ALT 343 12 130 U/LH
ALKP 315 14 111 U/LH Catalyst_One 07-04-22 12:02a GGT 12 0 4 U/LH TBIL 20 0 15 µmol/L

This submitted study contained 27 still images and 6 videos for review. Please submit primarily comprehensive videos.

ULTRASONOGRAPHIC EXAMINATION OF THE ABDOMEN

Urinary System

The urinary bladder, trigone, and cystourethral junction exhibited normal thickness and tone. Anechoic urine was present in the lumen with no uroliths or sediment. The ureteral papillae were normal. The ureters were not visible which is normal. No evidence of inflammatory or neoplastic changes was noted.

The area of the aortic trifurcation was free of pathology.

Normal size and margination were present in the kidneys. A normal 1:3 cortex / medulla ratio was maintained. The medulla and cortices were uniform in texture with some increased echogenicity and minor loss of corticomedullary symmetry and definition expected for the age of the patient. No evidence of pelvic dilation was present. The left kidney measured 3.9 cm in length. The right kidney measured 3.6 cm in length.

Adrenal Glands

The bilateral adrenal glands were overtly normal in size, position and shape. The left adrenal gland measured 0.30 cm width. The right adrenal gland measured 0.37 cm width.

Spleen

The spleen exhibited a finely textured and homogenous parenchyma which was hyperechoic to the liver and renal cortical parenchyma. The capsule was smooth and regular without apparent expansion. The splenic vasculature at the hilus was normal in volume with no evidence of congestion or thrombosis. Acute to chronic inflammatory, neoplastic, or benign parenchyma changes were not noted. The spleen measured 0.58 cm width at the level of the hilus.

Liver/ Gallbladder

The liver exhibited potential for subjective mild enlargement yet maintained symmetrical yet mildly rounded hepatic contour. Generalized uniform mild increased hepatic parenchyma echogenicity



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

compared to the spleen was present. No overt hepatic masses or nodules were noted. The gallbladder was non-distended in size. The gallbladder walls were sonographically normal without evidence of inflammatory criteria. Anechoic content was present primarily with minor mildly hyperechoic luminal debris. The area of the common bile duct was overtly normal without evidence of common bile duct dilation, stasis, or post hepatic obstructive criteria.

Gastrointestinal

The stomach was not definitively visualized.

The visualized segments of small intestine presented intact wall layering and subjectively maintained a 1:3 muscularis/mucosa ratio. The small Intestinal wall width measured 0.22 cm. The ileocolic wall width measured 0.34 cm.

Normal visible colon wall layers were present with subjective semi-formed to soft feces consistent with history.

Pancreas

The pancreas was normal in size and contour with isoechoic to heterogeneous parenchyma compared to adjacent omentum. No signs of active inflammation or neoplasia.

Free Abdomen

No overt omental masses, significant lymphadenopathy or evidence of peritoneal free fluid was present.

ULTRASONOGRAPHIC FINDINGS

- Hepatopathy exhibiting uniform parenchyma hyperechogenicity
- Minor gallbladder debris - no evidence of post hepatic obstructive criteria
- Overtly normal visualized small bowel
- Mild age-related kidneys

INTERPRETATION OF THE FINDINGS & FURTHER RECOMMENDATIONS

The overall liver was nonspecific with potential considerations including metabolic / vacuolar / reactive / inflammatory hepatopathy, i.e., cholangiohepatitis. Potential for occult round cell neoplasia such as lymphoma, which may present in a similar sonographic manner, cannot be excluded. Assuming normal clotting status and using a 25-gauge needle, ultrasound-guided hepatic FNA for cytology is warranted.

A GI panel to include PLI/TLI/Cobalamin/Folate is suggested to assess for nonobvious gastrointestinal disease or pancreatitis, given the patient's clinical signs. Empirically, and pending additional diagnostics, continued therapy for cholangiohepatitis +/- potential Triad Disease may prove beneficial.



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

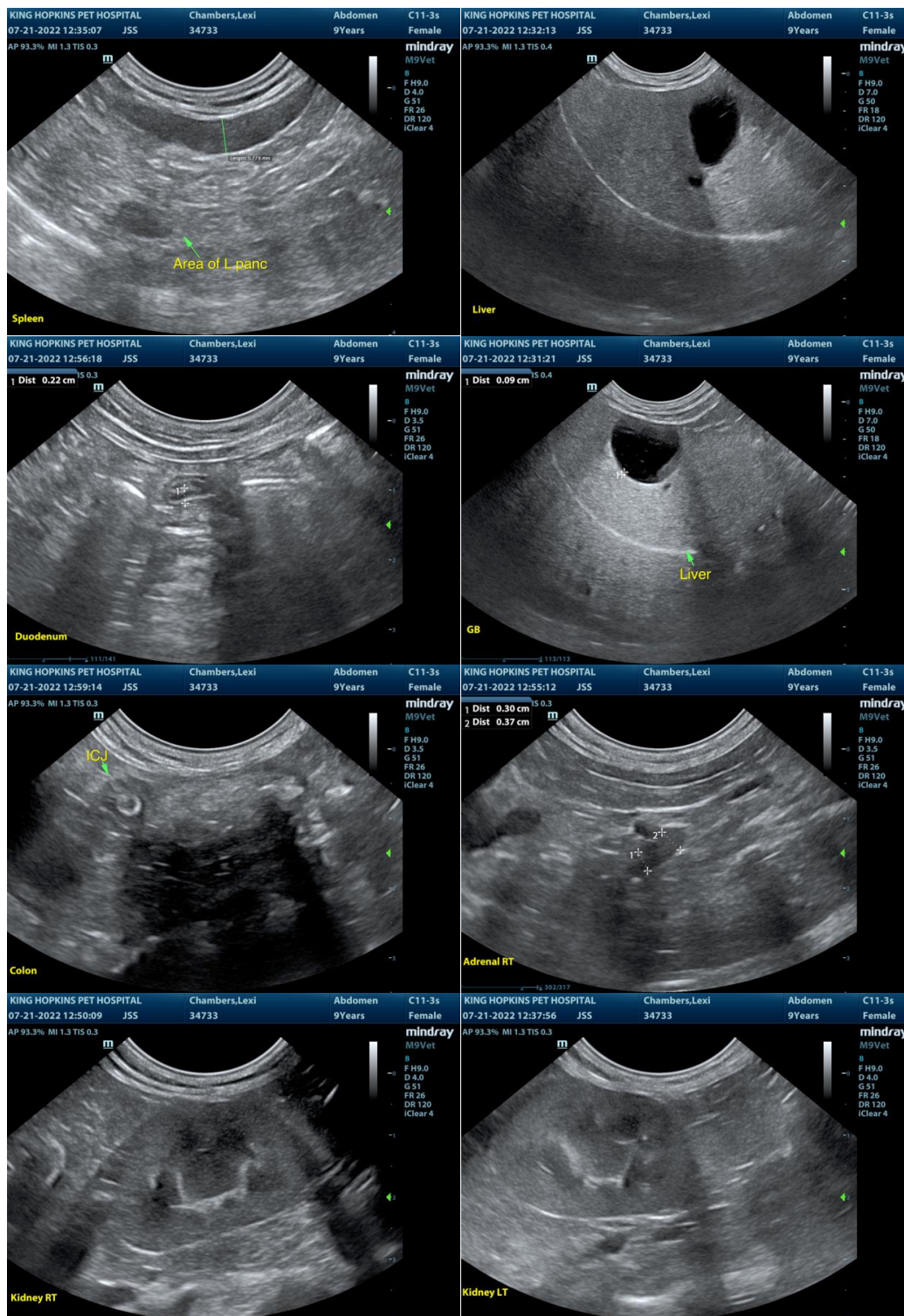
Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22





PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22



The information and recommendations provided are based on the images presented by the referring veterinarian/sonographer. No evaluation can be communicated regarding pathology that was not visible in the image/video clips provided.

Thank you for this referral. If the clinical or image interpretation does not parallel your findings or if I can be of any further assistance please contact me.

R. McKenzie Daniel, DVM, DABVP (Canine / Feline Practice)
info@SonoPath.com

Feline Liver Disease & Treatment Recommendations

<http://www.sonopath.com/FelineLiverDisease>

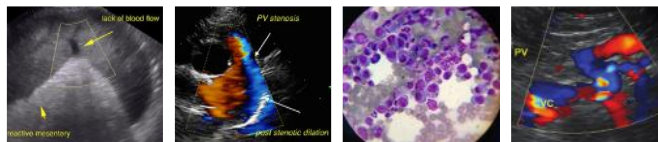
Description: Liver disease is a common clinical condition in cats; however, it can be subdivided into specific disease categories. Cats most often develop feline cholangitis, which is comprised of various forms of inflammatory liver disease or hepatic lipidosis; however, there are other disease processes, including neoplasia, infectious disease, and toxicities, that result in hepatic dysfunction. This chapter will focus predominantly on feline cholangitis and hepatic lipidosis; hepatic neoplasia is discussed in greater detail in a separate chapter.

1. Feline cholangitis complex is one of the most significant diseases in cats. The term “complex” embodies many different disease processes, each with its own signs and treatment protocols. The World Small Animal Veterinary Association (WSAVA) liver Standardization Group classifies feline cholangitis accordingly: neutrophilic cholangitis, which includes both acute and chronic neutrophilic cholangitis (these are likely extensions of the same disease process); lymphocytic cholangitis; and cholangitis associated with a liver fluke infection.

a) Acute neutrophilic cholangitis is a suppurative disease process of the liver and is most commonly seen in young to middle-aged cats. Acute cholangitis is almost always of bacterial origin, with enteric isolates being the most common culprits (these are thought to ascend from the biliary tree). Pancreatitis and inflammatory bowel disease (IBD) are associated disease processes. Histopathologically, this disease is represented by suppurative inflammation within the walls and



PATIENT	
Lexi Chambers	lumen of the biliary ducts, and may extend into the portal triads and possibly into the hepatic parenchyma.
SPECIES	b) Chronic neutrophilic cholangitis is either neutrophilic or lymphoplasmacytic, and arises from the acute form. Inflammation is centered around the portal region and includes lymphocytes, plasma cells, and neutrophils. Inflammation can also extend into the surrounding parenchyma and is sometimes noted within the lumen of the bile duct. Current research is investigating the role of <i>Bartonella</i> species in the development of chronic cholangitis; other postulated etiologies include a <i>Helicobacter</i> infection and immune-mediated processes. Biliary hyperplasia occurs secondary to the chronicity of the disease, and fibrosis and/or cirrhosis represent the end-stage manifestation of these foregoing disease processes. Cirrhosis is a rare condition in the cat because most cats either succumb or undergo successful treatment prior to this stage. Cats with chronic neutrophilic cholangitis often have chronic pancreatitis and IBD (triaditis). These other diseases may be responsible for the immune-mediated destruction of the liver that is commonly seen with the chronic form. The fact that the bile and pancreatic ducts are anatomically close to one another in cats may be the reason for coinfection and explain why the diseases occur in tandem.
Feline	
BREED	
3.1 kg	
SEX	
Spayed	
AGE	
10 years	
WEIGHT	
3.1 kg	Severe lymphocytic cholangitis is defined as chronic inflammation of the biliary tract infiltrated by small lymphocytes. Cats with lymphocytic cholangitis are usually ill for months or years. This disease is more common in Europe than North America, and is manifested as a chronic inflammatory disease, which ultimately leads to fibrosis and cirrhosis. Chronic infections are thought to be the result of <i>Helicobacter pylori</i> infections and immune-mediated diseases. There is a greater predisposition in both Persian and younger cats to chronic forms.
INTERPRETED BY	
R. McKenzie Daniel, DVM, DABVP (Canine and Feline)	c) Lymphocytic portal hepatitis is characterized by mild lymphocytic inflammation around the portal areas, but no inflammation within the bile ducts or hepatic parenchyma. This is a common finding upon biopsy, and although it can be nonspecific and incidental, it is also thought to indicate a reactive hepatopathy secondary to extrahepatic disease.
IMAGING PERFORMED BY	
JSS	
HOSPITAL NAME	d) In tropical and subtropical geographic areas, cholangitis is associated with fluke infestation secondary to infection with <i>Platynosomum</i> spp. Cats become infected by ingesting the second intermediate host (reptiles and amphibians). The fluke infection results in cystic dilation and bile duct thickening as well as obstruction.
King Hopkins PH	
REFERRING VET	
Dr. Latoya Brown	2. Hepatic lipidosis (HL) is one of the most common causes of hepatic disease in cats. It is defined as an accumulation of lipid within the cytoplasm of the hepatocyte. This can be idiopathic (primary) or can occur secondary to other diseases, and results in anorexia and weight loss. The pathophysiology is multifactorial and due to dysregulation of lipid metabolism in a catabolic state, which leads to excess accumulation of intracellular lipid within the hepatocyte.
INVOICE	
14347	
DATE	Other common causes of liver enzyme elevation in cats include hepatocellular thyrotoxicosis, infectious diseases (e.g. <i>Toxoplasma</i> and FIP), and primary and metastatic neoplasia. In hepatic toxicosis, thyroid hormones have a direct toxic effect on liver cells and stimulate increased liver enzyme activity. Moreover, increased intestinal motility secondary to hyperthyroidism can cause increased oxygen utilization and thus hepatic hypoxia, which ultimately leads to hepatic dysfunction. Acute cholangitis cannot be fully ruled out without conducting a biopsy. Owner (and patient)
7/21/22	



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

compliance with methimazole should be evaluated. The methimazole dose may need to be increased or other treatment modalities explored, including radioactive iodine, thyroidectomy, and/or dietary therapy with restricted iodine, such as the therapeutic diet, Hill's® y/d®. It should be noted that methimazole can also cause a drug-induced toxicosis and that this can also result in liver enzyme elevation.

Clinical signs:

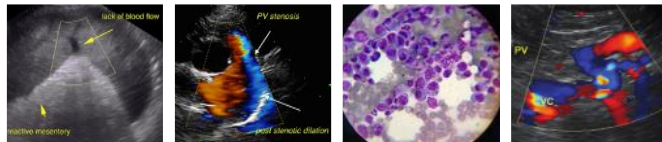
Cholangitis: Because of the acute suppurative nature of acute neutrophilic cholangitis, clinical signs commonly include pyrexia and jaundice, accompanied by vomiting, diarrhea, and lethargy. Of the four main complex types discussed above, acute cholangitis patients normally present as the most severely ill. A chemistry panel often reveals a moderate increase in ALT, ALP, GGT, bilirubin, and bile acids, while a CBC commonly shows an elevated white blood cell count (WBC) with or without a left shift.

Cats suffering from the remaining types of feline cholangitis normally display less severe signs, but have likely been sick for a longer period of time. Those with chronic neutrophilic cholangitis can have intermittent episodes of jaundice and vomiting, which are cyclic and self-resolving. Weight loss, anorexia, and lethargy are common, and one typically observes elevations in ALP, GGT, bilirubin, and bile acids. The degree of elevation in ALT and AST is variable.

Patients with severe lymphocytic cholangitis exhibit weight loss and anorexia; however, because it is a slow-moving, progressive disease, signs may be chronic and mild. Liver enzyme elevations are generally mild until the chronic phase when icterus occurs as well as ascites. Hypergammaglobulinemia is also a prominent feature of this disease.

Cats with fluke infestations are either asymptomatic or systemically ill with pyrexia, vomiting, anorexia, icterus, and bile duct obstruction. A CBC may indicate marked liver enzyme elevations as well as an eosinophilia. Cats with lymphocytic portal hepatitis are asymptomatic and do not demonstrate laboratory abnormalities.

Hepatic lipidosis: The most common clinical signs are anorexia, vomiting, diarrhea, icterus, lethargy, depression, ptyalism, and weight loss. Since HL occurs in the face of so many other diseases, clinical signs may vary and be due, in part, to the underlying disease process. Likewise, lab work abnormalities can also vary, depending on concurrent disease processes. Hepatic encephalopathy may ensue, resulting in severe weakness, depression, and ptyalism. Common CBC abnormalities include a nonregenerative anemia, stress leucogram, poikilocytosis, and the presence of Heinz bodies. On the serum chemistry, an elevation in ALP is disproportionate to GGT levels, which are usually within normal limits. Serum ALT is variably increased, but typically of a lower magnitude than that of ALP. Bilirubin is increased due to intrahepatic cholestasis. Bile acids are increased, but are a superfluous indicator in the face of hyperbilirubinemia. The BUN and albumin may be normal or subnormal. Coagulation abnormalities occur in cats with HL due to vitamin K deficiency, which is a result of malabsorption in the intestines and decreased production of coagulation factors in the liver due to severe hepatic dysfunction.



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

Cats with thyroid toxicosis mainly exhibit clinical signs of hyperthyroidism. The chief abnormalities on serum chemistry are a mild to moderate elevation in ALT and elevated T4 levels. In the event of methimazole toxicity, the ALT can also be mild to moderately elevated. Cats with hepatic neoplasia may also demonstrate similar signs to other forms of hepatic disease, display elevations in ALT, ALP, GGT, and AST, and exhibit a leukocytosis and possibly anemia of chronic disease. This condition is discussed in greater detail in the chapter on "Hepatic Neoplasia."

Diagnostics: Ultrasound and interventional ultrasound are important means of definitively diagnosing hepatic disease in cats. An ultrasound-guided core biopsy of the liver can be performed to acquire both histopathology and aerobic/anaerobic cultures. Alternatively, a larger tissue biopsy can be obtained via laparoscopy or laparotomy. In the event that a laparotomy or laparoscopy is performed, biopsies of the small intestine and pancreas should be obtained. One will also often encounter triaditis. Ultrasound-guided centesis of the gallbladder can be done to collect bile for culture and is preferred over hepatic parenchymal cultures. Ultrasound additionally allows for visualization of choleliths and obstruction to bile flow; it also helps determine whether surgical intervention is required. In the case of HL, the sonographic appearance presents as a uniform, diffuse, dense hyperechoic parenchyma that is hyperechoic to falciform fat and spleen; however, a primary underlying disease may also be present, such as cholangitis or lymphoma. In cats with suspected HL, fine needle aspiration (FNA) is the safest way to rule this out and to evaluate for lymphoma, as cytology is especially useful in the diagnosis of both these diseases. Lipidotic livers are friable and hence bleeding can occur as a complication of biopsy due to poor tissue integrity, lack of tissue hemostasis, and possibly compromised systemic hemostasis due to poor hepatic function. Cholangitis and neoplasia can be suspected on cytology, but a hepatic biopsy is preferred in order to define architecture, inflammatory infiltrate, and fibrosis. The clinician must weigh the risks and benefits of obtaining a biopsy in patients that may have a concurrent disease or are unresponsive to conventional therapy for HL. Pretreatment with vitamin K1 and aggressive supportive care may aid in stabilizing a patient for biopsy.

Cholangitis associated with liver flukes is uncommon in North America, but if the patient is in a tropical or subtropical location, the diagnosis is either obtained by fecal examination or liver biopsy, which permits observation of the flukes and/or their eggs within the bile ducts.

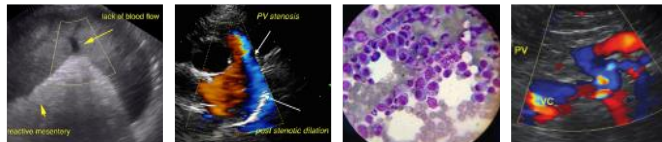
Treatment: The following medications are suggested in keeping with general guidelines for treatment for feline hepatic disease; however, each patient should be assessed and treated as an individual, and management should be tailored according to a specific diagnosis.

Disease-specific recommendations:

1. Feline cholangitis complex

a) Acute neutrophilic cholangitis and chronic neutrophilic cholangitis:

- Antibiotics: Administration of antibiotics should ideally be based on culture and sensitivity. If a culture is not available, one may treat with broad-spectrum antibiotics, such as amoxicillin (10-20 mg/kg PO BID), amoxicillin clavulanic acid (10 mg/kg PO BID or 62.5 mg PO BID), a cephalosporin, such as cefadroxil (20 mg/kg PO BID), or enrofloxacin (5 mg/kg



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

PO once daily). Antibiotics can be used for 4-8 weeks. Metronidazole (11-22 mg/kg PO BID) can be given as an anti-inflammatory medication; it also has an anaerobic spectrum. Metronidazole also decreases ammonia produced by intestinal microbes (administer at 7.5 mg/kg PO BID-TID in cases of hepatic encephalopathy).

b) Chronic neutrophilic cholangitis (lymphoplasmacytic inflammation):

- An anti-inflammatory medication should be given when biopsy reveals that there has been significant infiltration with lymphocytes and/or plasma cells, or if the patient is not responding to antibiotic medication alone (in the absence of a biopsy).
- Prednisolone should be given at 1-2 mg/kg/day. Start at a higher dose, and wean over time to every other day in decreasing dosages every 2-4 weeks following the resolution of signs. Additional immunosuppressant medications are not typically used in cats for this disease.

c) Severe lymphocytic cholangitis:

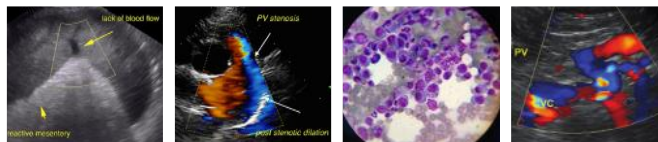
- Prednisolone should be dosed at 1-2 mg/kg/day; however, it remains controversial as to whether prednisolone is in fact effective in the course of this disease.
- Ursodiol (Actigall): 10-15 mg/kg PO Q24hr.

d) Liver fluke infection:

- Praziquantel: Give 20-30 mg/kg PO Q24hr for 3 days.

2. Hepatic Lipidosis

- Hyperalimentation is crucial in the management of HL; it may also be necessary in the management of other hepatic diseases in cats that are not eating so as to prevent hepatic lipidosis as a complication. In order to determine the caloric needs of the patient, calculate the basal energy requirement using the formula $BER = 70 \times BW \text{ kg}^{0.75}$. Multiply the BER by an illness energy requirement factor (1.25-1.4 in cats) and then select a therapeutic recovery diet with enhanced protein and fat levels, such as Hill's® a/d®, CliniCare® Liquid Diet (Abbott Animal Health), Royal Canin® Recovery RS™, or Iams® Maximum-Calorie™. Feed small, frequent meals through an esophagostomy tube (E tube), percutaneous endoscopic gastrostomy tube (PEG), or nasoesophageal tube (NE) tube. Give slowly over 15-30 minutes or trickle feed as a CRI. In cats that have been anorexic for a prolonged period of time, the amount of food should be gradually increased over 3 days' time. The food should be made into a slurry and warmed, and the total amount of food divided into 4-6 feedings per day. Flush the feeding tube with 5-15 ml warm water. Pretreatment with cisapride (1.25-2.5 mg/cat PO or 0.1-0.5 mg/kg PO BID-TID) or metoclopramide (0.2-0.4 mg/kg PO or SC Q8hr) can prove helpful to improve gastric emptying; dosing should occur 30 minutes before feeding.
- Vitamin K1: Give 0.5-1.5mg/kg SQ or IM every 12 hours for a maximum of 2-3 doses if clotting times are increased. (The latter commonly occurs in the face of hepatic lipidosis due to decreased intestinal absorption of vitamin K as well as hepatic failure.)



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

- L-carnitine: Give 50-100 mg/kg PO Q24hr. L-carnitine is indicated in cats with severe hepatic lipidosis.
- Taurine: Give 250-500 mg PO Q24hr. Taurine can be administered as a supplement; it is an essential amino acid in cats.

3. Hepatic Encephalopathy:

- Lactulose: Give 0.5 ml/kg PO BID-TID to soften the stool. Lactulose helps manage hepatic encephalopathy by combining with ammonium in the GI tract and thus decreasing circulating ammonia levels. It can also be mixed into the slurry during feeding. Lactulose can also be given as a retention enema in an encephalopathic crisis.
- Metronidazole or neomycin: Give metronidazole at 7.5 mg/kg PO BID-TID. This is an antimicrobial, which reduces bacterial counts and reduces ammonia production in the colon. Alternatively, administer neomycin at 20 mg/kg PO BID-TID.
- L-Carnitine: Give 50-100 mg/kg PO Q24hr. Normally synthesized by the liver, L-carnitine enhances ammonia elimination and is indicated in cases of hepatic encephalopathy and lipidosis. Carnitine must be in the L-form.
- Diet: A low-protein diet with high amounts of biologically available protein is recommended in encephalopathic patients to reduce the nitrogen load from the breakdown of amino acids.

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

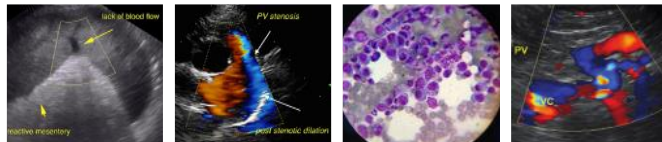
14347

DATE

7/21/22

General treatment recommendations for cats with either feline cholangitis complex or hepatic lipidosis:

- IV Fluids: Fluid therapy is integral, especially in cats with severe liver disease as they are often inappetant and dehydrated. In the face of hepatic failure, avoid Lactated Ringer's solution (LRS) as lactate is metabolized by the liver. Monitor electrolytes closely. Add potassium in the form of potassium chloride (KCL); 20 mEq/L is a general starting point, but higher doses may be needed in the face of severe hypokalemia (maximum infusion rate is 0.5 mEq/kg/hr). Correct concurrent hypomagnesemia with magnesium sulfate or magnesium chloride at 0.75-1 mEq/kg/day given as a CRI for one day, and then reduce it to 0.3-0.5 mEq/kg/day. Monitor serum phosphorus levels and supplement as needed. Hypophosphatemia can occur following the reinstatement of feeding, especially in previously anorexic patients (re-feeding syndrome). Supplement phosphorus at 0.01-0.06 mmol/kg/hr using potassium phosphate or sodium phosphate.
- Vitamin B1 complex (thiamine) can also be added to the fluids at 1-2ml/liter. Note: Protect the fluid bag from light as the vitamins degrade when exposed to light.
- Vitamin B12 (cyanocobalamin) can be administered at 250 ug SC or IM weekly as needed in cases of HL or in cats with primary gastrointestinal disease.
- Famotidine can be given 0.5 mg/kg PO or IV once to twice daily as an antacid.
- Ursodiol (Actigall): Give 10-15 mg/kg PO Q24hr, with food, to stimulate bile secretion and flow, and decrease cholestasis. It has immunomodulatory, anti-fibrotic, and choleric effects and anti-copper storage benefits; it also stabilizes mitochondrial function. Ursodiol can be compounded into a liquid formulation for cats.
- S-adenosylmethionine (SAME): Give 90 mg/cat PO on an empty stomach (1-2 hours before feeding), or a loading dose of 35-60 mg/kg once to twice daily and a maintenance dose of



PATIENT

Lexi Chambers

SPECIES

Feline

BREED

3.1 kg

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

IMAGING PERFORMED BY

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22

20 mg/kg PO Q24hr. SAMe replenishes glutathione and aids in cellular detoxification. It is also an anti-inflammatory and antioxidant.

- Antiemetics: these are used to decrease the frequency of vomiting and therefore enable enteral nutrition. A common antiemetic, metoclopramide (0.2-0.5 mg/kg SC TID 30 min before feeding or 0.01-0.02 mg/kg/hr as a CRI) has the beneficial effect of concurrently improving gastric emptying. Alternative anti-emetics include: maropitant citrate (Cerenia), which should be administered at 1 mg/kg subcutaneously once daily for up to 5 days; odansetron (Zofran), which can be dosed at 0.1 mg/kg PO once to twice daily or 0.1-0.3 mg/kg IV BID-TID; or dolasetron (Anzemet), which can be administered at 0.5 mg/kg PO, SC, or IV Q24hr. Silybin-phosphatidylcholine (Marin) (5 mg/kg PO Q24hr) is yet another alternative; however, to date there are no evidenced-based studies in cats on the effects of milk thistle. Nevertheless, it is suggested that it acts as an antioxidant and free radical scavenger, decreases hepatotoxin binding, improves glutathione concentrations, aids in iron chelating, and promotes choleresis.
- Vitamin E: Give 10-15 IU/kg/day PO (100-400 IU) in a water-soluble form twice daily.

Initially, in the face of hepatic failure and severe icterus, patients are hospitalized and monitored closely. Once stable, the patient can be discharged with a feeding tube if oral intake is not yet sufficient. Weight, appetite, and blood work must be carefully assessed. Specifically, ALT, SAP, serum protein, and serum albumin should initially be evaluated every 2 weeks, then monthly for the first 6 months, and subsequently every 4-6 months. A patient's response to therapy is typically assessed by their clinical condition and laboratory work; however, repeated biopsy or FNA may be necessary if the clinical response proves to be unsatisfactory.

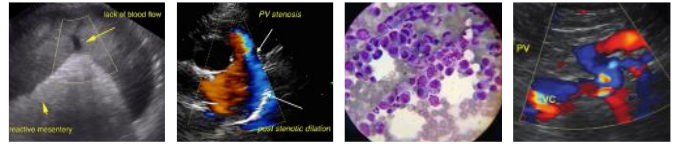
Common drugs to avoid when treating hepatic disease: Halothane; sulphonamides; diazepam; azole antifungals; phenobarbital; tetracyclines; erythromycin or enrofloxacin if combined with theophylline or cisapride; and cimetidine if combined with theophylline, metronidazole, or chloramphenicol. Sedate with caution if lipidosis is present. If anesthesia is necessary for sampling or placing a feeding tube, induction with propofol should be considered with an inhalant, such as sevoflurane or isoflurane.

Conclusion: There are many pathological processes that can result in hepatic disease in the feline patient. Ideally, sampling is performed to obtain a definitive diagnosis, which allows for specific guidance with regards to therapy and prognosis. Adequate nutrition or hyperalimentation is critical for hepatic support when treating HL or preventing it from occurring as a secondary complication. Intensive care and monitoring are also key components of therapy in critically ill icteric cats.

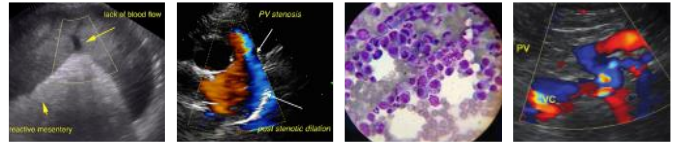
References:

Armstrong PJ, Blanchard G. Hepatic lipidosis in cats. *Vet Clin North Am Small Anim Pract* 2009;39(3):599-616.

Brian PH, Barrs VR, Martin P, et al. Feline cholecystitis and acute neutrophilic cholangitis: clinical findings, bacterial isolates and response to treatment in six cases. *J Feline Med Surg* 2006;8(2):91-103.



PATIENT	Brenner K, KuKanich KS, Smee NM. Refeeding syndrome in cat with hepatic lipidosis. <i>J Feline Med Surg</i> 2011;18(8):614-17.
Lexi Chambers	
SPECIES	Carreira VS, Vieira RF, Machado GF, Luvizotto MC. Feline cholangitis, cholangiohepatitis complex secondary to <i>Platynosomum fastosum</i> infection in a cat. <i>Rev Bras Parasitol Vet</i> 2008;17 Suppl 1:184-87.
Feline	
BREED	Center S. Feline hepatic lipidosis is a common liver disorder. <i>Feline Health Topics</i> 1989;4(3):1-6.
3.1 kg	
SEX	Center SA, Baldwin BH, Dillingham S, et al. Diagnostic value of serum gamma-glutamyl transferase and alkaline phosphatase activities in hepatobiliary disease in the cat. <i>J Am Vet Med Assoc</i> 1986;188:507-10.
Spayed	
AGE	Davenport D. Antimicrobial therapy for gastrointestinal, pancreatic, and hepatic disorders. <i>Probl Vet Med</i> 1990;2(2):374-93.
10 years	
WEIGHT	Gagne JM, Armstrong PJ, Weiss DJ, et al. Clinical features of inflammatory liver diseases in cats: 41 cases (1983-103). <i>J AM Vet Med Assoc</i> 1999;214:513.
3.1 kg	
INTERPRETED BY	Greiter-Wilke A, Scanziani E, Soldati S, et al. Association of <i>Helicobacter</i> with cholangiohepatitis in cats. <i>J Vet Intern Med</i> 2006;20(4):822-27.
R. McKenzie Daniel, DVM, DABVP (Canine and Feline)	
IMAGING PERFORMED BY	Holan KM. Feline Hepatic Lipidosis. In: Bonagura JD, Twedt DC, eds: <i>Current Veterinary Therapy XIV</i> . St. Louis, MI: Saunders Elsevier; 2009:570-75.
JSS	
HOSPITAL NAME	Kordick DI, Brown TT, Shin K, Breitschweidt EB. Clinical and pathologic evaluation of chronic <i>Bartonella henselae</i> or <i>Bartonella clarridgeiae</i> infection in cats. <i>J Clin Microbiol</i> 1999;37(5):1536-47.
King Hopkins PH	
REFERRING VET	Mathews KA. Nutritional Support. In: Mathews KA, ed: <i>Veterinary Emergency and Critical Care</i> . Guelph, ON: Lifelearn Inc, 1996:19.1-19.29.
Dr. Latoya Brown	
INVOICE	Otte CM, Penning LC, Rothuizen J, Favier RP. Retrospective comparison of prednisolone and ursodeoxycholic acid for the treatment of feline lymphocytic cholangitis. <i>Vet J</i> 2013;195(2):205-9.
14347	
DATE	Tams T. Management of liver disease in dogs and cats. <i>Mod Vet Pract</i> 1984;65(2):107-14.
7/21/22	
	Twedt DC, Armstrong PJ. Feline inflammatory liver disease. In: Bonagura JD, Twedt DC, eds: <i>Current Veterinary Therapy XIV</i> . St. Louis, MI: Saunders Elsevier; 2009:576-581.
	van den Ingh TSGAM, Cullen JM, Twedt DC, et al. Morphological classification of biliary disorders of the canine and feline liver. In: Rothuizen J et al, eds: <i>WSAVA Standards for Clinical</i>



PATIENT

Lexi Chambers

and Histological Diagnosis of Canine and Feline Liver diseases. Spain: Saunders Elsevier; 2006:61-78.

SPECIES

Feline

Webster C, Cooper J. **Therapeutic** use of cytoprotective agents in canine and feline hepatobiliary **disease.** *Vet Clin North Am Small Anim Pract* 2009; 39(3):631-52

BREED

3.1 kg

Weiss D, Armstrong P, Gagne J. **Inflammatory liver disease.** *Semin Vet Med Surg (Small Anim)* 1997;12(1):22-27.

SEX

Spayed

AGE

10 years

WEIGHT

3.1 kg

INTERPRETED BY

R. McKenzie Daniel,
DVM, DABVP
(Canine and Feline)

**IMAGING
PERFORMED BY**

JSS

HOSPITAL NAME

King Hopkins PH

REFERRING VET

Dr. Latoya Brown

INVOICE

14347

DATE

7/21/22