


PATIENT

Henry Freylikham

PRESENTING CLINICAL SIGNS

Weight loss, possible opacity mid abdomen. Newly acquired heart murmur.

SPECIES

Canine

BREED

Coton de Tulear

SEX

Neutered Male

AGE

2008

WEIGHT

11.8 Pounds

ULTRASONOGRAPHIC EXAMINATION OF THE HEART & ABDOMEN

CANINE CARDIAC PARAMETERS	MR VMAX (m/s)	TR VMAX (m/s)	LA/AO (Boon method)	LA/AO (Heart Base; Swe)	FS (%)	EF (%)	EPSS (cm)
NORMAL PARAMETER	4.5-5.5	<2.7	1.3	<1.6	28-40	40-100	<0.6
PATIENT	6.81		1.3	1.25	49	83	0.1
CANINE CARDIAC PARAMETERS	HR (BPM)	AV VMAX (m/s)	PV MAX (m/s)	BODY WEIGHT (kg)	LA 2D short axis Base view (cm)	LVIDd Avg; 2D and m-mode short axis (cm)	LVIDs Avg; 2D and m-mode short axis (cm)
NORMAL PARAMETER	50-100	0.7-1.7	0.7-1.6	BELOW	BELOW	BELOW	BELOW
PATIENT	182	1.48	0.74		2.87	2.27	

Cardiac Presentation

The echocardiogram in this patient demonstrated normal **left atrial** size based on 3 different LA measurement methods. Chamber volumes and echogenicity were normal. The cranial and caudal **mitral** valve leaflets presented vegetative thickening consistent with endocardiosis. Doppler indicated measurable insufficiency. The **left ventricle** presented thicknesses with linear contour and was not dilated nor restricted. The **myocardium** presented normal echogenicity without subjective evidence of significant fibrotic or ischemic disease. **Contractility** of the ventricular walls was adequate and in normal range for this patient evidenced by the fractional shortening measurement and subjective evaluation of the different regions of the myocardium. The **left ventricular outflow** tract demonstrated normal laminar flow and subjective structural integrity. The **right atrium** and auricle revealed normal size, structure and content. No evidence of masses was noted or chamber overload. **Tricuspid** valvular assessment demonstrated adequate linear morphology. The **right ventricle** was of normal size (1/3 diameter of LV), chordae structure, myocardial echogenicity and thickness. **Pulmonic** tract assessment revealed normal valve structure, laminar flow, and diameter (approx. 1:1 pa/ao ratio). No visible **pericardial** or free pleura fluid was noted. No echographically detectable evidence of infiltrative disease was visible. The cranial **mediastinum** and **pericardial** regions were free of masses in the visible window.

Urinary System

The **urinary bladder**, trigone, and pelvic urethra presented normal thicknesses and normal tone. The ureters were not visible which is normal. No uroliths or sediment were visualized and anechoic urine was present. No evidence of inflammatory or neoplastic changes were noted. Ureteral papillae were normal.

The **kidneys** revealed largely normal size and structure, corticomedullary definition and ratio (cortex 1/3 of medulla) were essentially maintained with some age-related loss of curvilinear patterns regarding the capsule and C/M junction. The cortices presented largely uniform texture with some increased echogenicity expected for his age patient. Medullary structure differed distinctly from that of the cortex

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and no evidence of pelvic dilation was present. The right kidney measured 4.47 cm. The left kidney measured 3.5 cm.

Adrenal Glands

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The **right adrenal gland** presented normal size and contour, measuring 1.73 cm x 0.65 cm.

The **left adrenal gland** was slightly enlarged at the cranial pole, measuring 1.05 cm at the cranial pole and 0.91 cm at the caudal pole, 2.16 cm in length. Increased in size approximately 50% more than normal.

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Spleen

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The **spleen** presented a smooth homogeneous parenchyma hyperechoic to liver and renal cortical parenchyma. The capsule was smooth without noticeable expansion or deviation from within the spleen or adjacent pathology. The splenic vasculature demonstrated normal volume without signs of congestion or thrombosis. No sonographic evidence of acute or chronic inflammatory, neoplastic, or infarctual changes were noted.

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Liver

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The **liver** images submitted revealed subjectively normal liver size, contour, and structure. Parenchymal echogenicity was naturally coarse and hypoechoic to the spleen. Vascular and biliary tracts were of normal volume with no evidence of congestion. The gallbladder presented acceptably thin walls with primarily anechoic content. The cystic and common bile ducts were normal. No pathological hepatic lymphadenopathy was evident. No overt structural evidence of inflammatory, infiltrative or regenerative pathology was evident.

Gastrointestinal

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The **stomach** itself was unremarkable. A 4.87 cm x 5.36 cm jejunal mass was noted cranial to the urinary bladder with minor serosal escape into the regional omentum caudally towards the bladder itself. This mass is stricturing with a mild amount of obstructive pattern.

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Pancreas

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The base and limbs of the **pancreas** were observed to be largely isoechoic to surrounding omental fat. Pancreatic duct and capsular contour were acceptably normal and parenchyma respected normal curvilinear patterns. No overt evidence of active inflammatory or neoplastic disease was noted.

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ULTRASONOGRAPHIC FINDINGS

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- Stage B1 valvular disease
- Jejunal mass with minor capsular escape, potentially resectable – Suspect leiomyosarcoma, round cell neoplasia or carcinoma possible, granulomatous disease unlikely.
- Enlarged left adrenal gland – Pheochromocytoma, adenocarcinoma, or benign hyperplasia possible.

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INTERPRETATION OF THE FINDINGS & FURTHER RECOMMENDATIONS

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Blood pressure measurements recommended. If hypertension is present, then urine catecholamine would be indicated. Chest radiographs recommended for screening. FNA of the intestinal mass or direct exploratory surgery with resection and anastomosis of the intestinal mass and left adrenalectomy indicated.

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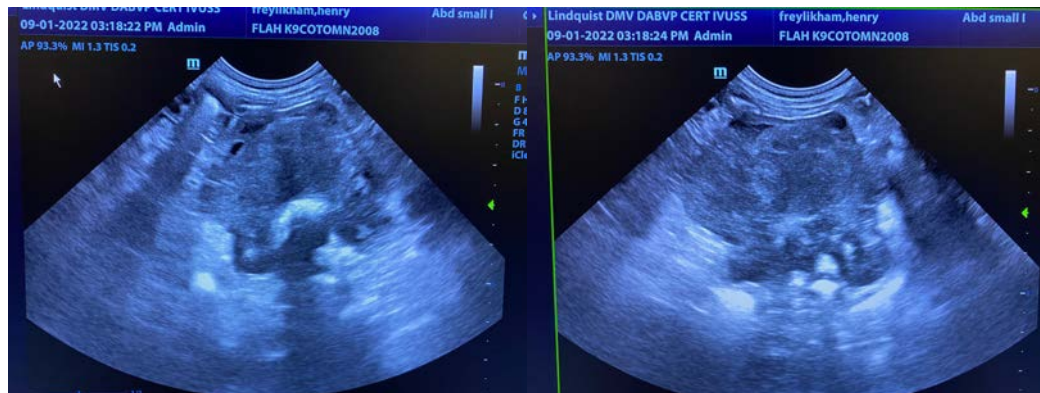
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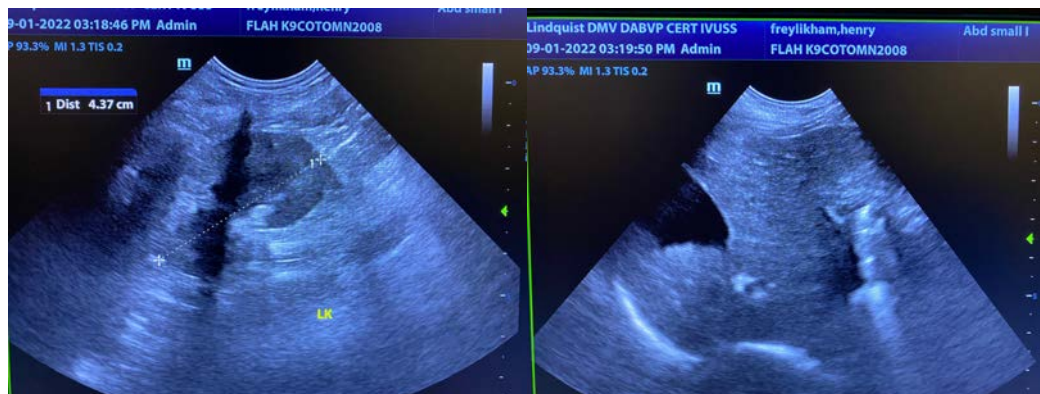
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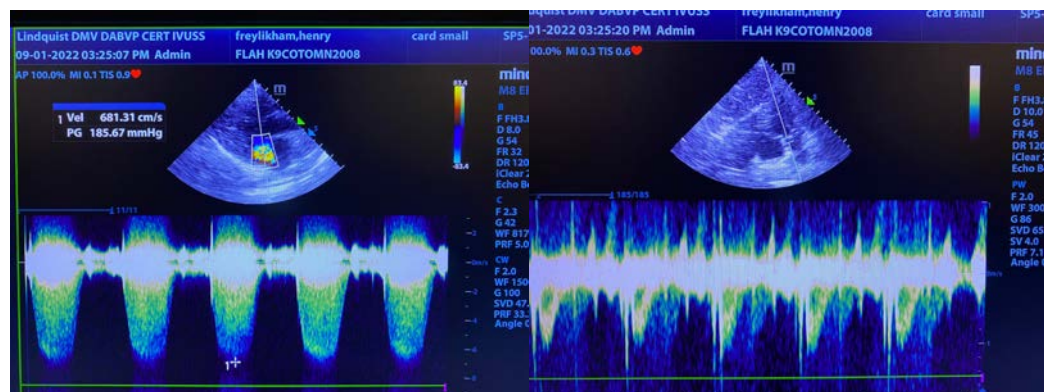
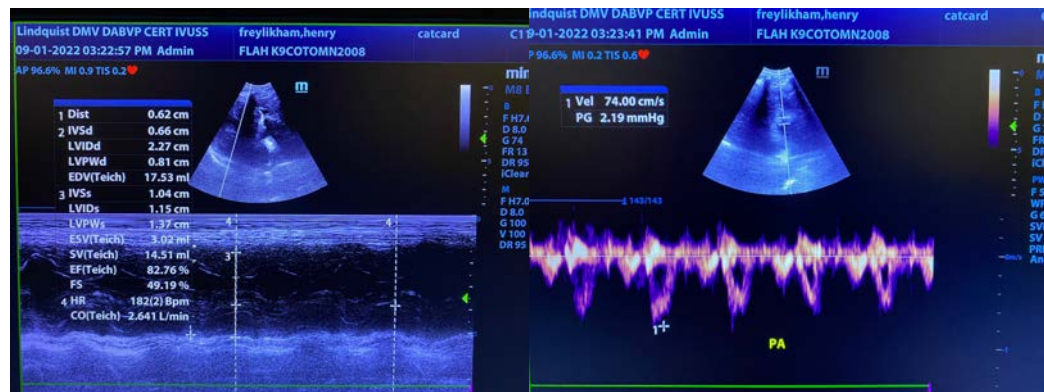
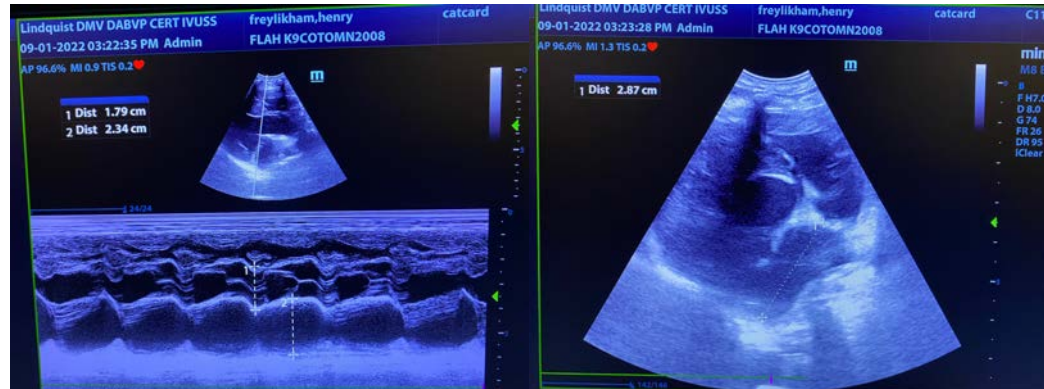
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The information and recommendations provided are based on the images presented by the referring veterinarian/sonographer. No evaluation can be communicated regarding pathology that was not visible in the image/video clips provided.

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.

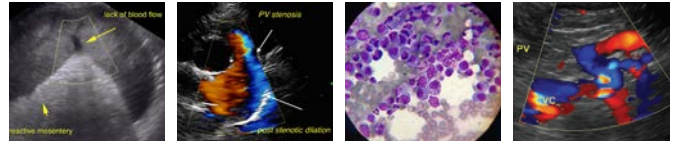
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Mitral & Tricuspid Valve Disease

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

Description: Myxomatous valvular degeneration refers to a sterile degenerative disease that affects middle-aged and older dogs. The mitral valve apparatus includes the left ventricular papillary muscles, chordae tendinae, valvular annulus, and anterior and posterior leaflets. The anterior leaflet is continuous with the aortic outflow tract. The accumulation of mucopolysaccharides within the spongiosa and fibrosa layers of the leaflets creates a vegetative nodular appearance. This vegetative pathology may also be the result of bacteremia due to dental disease or a different source of bacterial pervasion. Lengthening of the chordae tendinae occurs secondary to any excessive turbulent forces that might transpire. Chondrodystrophic breeds are overrepresented and typically have collapsing trachea and intervertebral disc disease. Cavalier King Charles Spaniels are paradigmatic of this phenomenon; they develop mitral valve disease (MVD) at an early age. MVD prevalence can be as high as 33% in toy breeds over 10 years of age and is the most common cardiac disease in dogs (it accounts for 75-80% of all cases). MVD has also been reported in German Shepherds; they demonstrate a more rapid progression of the disease in comparison to other breeds.



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Distortion of the mitral valve leaflets prevents normal coaptation of blood resulting in regurgitative flow of the stroke volume into the left atrium. The lengthening and rupture of the chordae can lead to leaflet prolapse into the atrium and the eventual inversion of the flail leaflet into the atrium, which will in turn exacerbate the regurgitation. Chordae rupture often occurs suddenly, leading to the rapid onset of congestive heart failure (CHF), as the myocardium does not have adequate time to hypertrophy and compensate for the defect. (A small amount of regurgitation can be well tolerated if it progresses more slowly over time.) The result is volume overload of the left atrium and left ventricle, which gives rise to eccentric hypertrophy. Eccentric hypertrophy causes dilation of the mitral annulus, which further complicates the myocardial left atrium and left ventricle stretch. Myocardial oxygen deprivation due to poor coronary perfusion (stretch and catecholamine stimulation) leads to myocardial cell death, replacement fibrosis, and myocardial dysfunction. This pathological progression is known as “overload cardiomyopathy.”

Clinical Signs: The most common sign of MVD is a progressive cough; it is due to left-sided volume overload, which in turn precipitates pulmonary edema and left atrial bulging at the left mainstem bronchus. Respiratory distress ensues gradually in chronic cases or rapidly in cases of primary chordae rupture. Exercise intolerance, syncope, ascites, weight loss, and anorexia are also commonly observed. A physical exam will often reveal an audible murmur best heard over the left cardiac apex. The murmur audibility and length into systole typically corresponds to the degree of mitral regurgitation but not necessarily to the severity of the cardiac status. Concurrent systemic hypertension may induce progression of the disease thereby necessitating systemic blood pressure measurements during medical management.

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A challenging situation occurs when the practitioner must determine whether the origin of a cough is respiratory (e.g. tracheal collapse, chronic bronchitis, COPD) or cardiac (e.g. pulmonary edema, mainstem bronchus pressure). A cough history of months or years, normal to high body scores, and normal heart rates with sinus arrhythmia tend to support chronic airway disease. In cases of MVD, however, the disease tends to be progressive: the body score is lower, the heart rate is higher, and there is a possibility of pathological arrhythmias. Yet, both bronchial disease and MVD can be present simultaneously, and advanced diagnostics, such as ultrasound examination, are necessary to distinguish which culprit is inciting the clinical signs.

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Diagnostics: Ascertaining NT-proBNP levels may help to determine whether dyspnea is respiratory or cardiac in origin and provide information that can assist with the early detection of cardiac disease in subclinical patients, especially in high-risk breeds, such as Cavalier King Charles Spaniels, Podles, and Cocker Spaniels.

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Radiographic findings may reveal a vertical or bulging caudal cardiac waist, hilar edema, generalized cardiomegaly, elevation of the left mainstem bronchus, and right-sided enlargement in advanced cases where pulmonary hypertension or tricuspid disease is also present.

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Echocardiographic findings allow the practitioner to assess precisely cardiac function under pathological circumstances. Moreover, they enable an evaluation of the myocardial response at the time of the exam and help practitioners continue to quantify that response over time while patients are undergoing therapy.

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Treatment: The following treatment options are based on the ACVIM consensus statement regarding MVD.

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Stage A: There is a high risk of cardiac disease, but as there are no clinical signs, no specific therapy—medical or dietary—is indicated. A radiograph and an echocardiogram should be conducted in one year or earlier if the patient is a large-breed dog as MVD may progress faster.

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Neutered Male

Stage B: Heart disease is present.

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B1: There is a murmur, but no chamber enlargement. Treatment is the same as for stage A.

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B2: There is a murmur with left atrial and left ventricular enlargement, but the patient is asymptomatic. Start angiotensin-converting enzyme (ACE) inhibitors (enalapril at 0.5 mg/kg PO Q12-24hr or benazepril 0.25-0.5 mg/kg PO Q24hr). Consider beta blockers and mild sodium restriction.

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Stage C: There are past or current clinical signs of heart failure.

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Acute CHF:

- Lasix 2 mg/kg IV or IM hourly to a total dose of 8 mg/kg until the respiratory rate has normalized. Alternatively, for cases of life-threatening pulmonary edema, administer as a CRI (1mg/kg/hour).
- Oxygen supplementation.
- Continue with the ACE inhibitor and add pimobendan at 0.25-0.3 mg/kg PO BID.

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Chronic therapy:

- Lasix at 2 mg/kg PO BID and increase incrementally as needed.
- Continue with the ACE inhibitor.
- Pimobendan at 0.25-0.3 mg/kg PO BID.

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- Consider adding spironolactone at 0.25-2 mg/kg PO BID to control congestion.
- A sodium-restricted diet is recommended, although some dogs will not eat it.

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- Consider administering omega-3 fatty acids, digoxin, theophylline, and cough suppressants.
- If the dog is not already on beta blockers, then do not commence.

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Stage D: This is the end-stage of the disease. Continue with the standard therapy of diuretics, ACE inhibitors, and pimobendan, and consider the following:

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Neutered Male

- Abdominocentesis when applicable to decrease discomfort if the patient is undergoing respiratory distress.
- Anti-anxiety medications for sedative purposes.
- Sodium nitroprusside and dobutamine (in a critical care facility).
- Nitroglycerin.

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We recommend monitoring serum urea, creatinine, electrolytes, urine specific gravity, and possibly blood pressure for 5-7 days after therapy has commenced. A repeat ECG is warranted if an arrhythmia was present during the original assessment.

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References:

Atkins C, Bonagura J, Ettinger S, et al. Guidelines for the diagnosis and treatment of canine chronic valvular heart disease. *J Vet Int Med* 2009; 23(6):1142-50.

Borgarelli M, Buchanan J. Historical review, epidemiology and natural history of degenerative mitral valve disease. *J Vet Cardiol* 2012;14:93-101.

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Fox P. Pathology of myxomatous mitral valve disease in the dog. *J Vet Cardiol* 2012;14:103-26.

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