



PATIENT

Remy Bartone

SPECIES

Canine

BREED

Yorkie

SEX

Neutered Male

AGE

13 Years

WEIGHT

9.6 kg

INTERPRETED BY

Eric Lindquist, DMV

DABVP, Cert. IVUSS

IMAGING PERFORMED BY

Erin Wicks

HOSPITAL NAME

Shores VEC

REFERRING VET

Dr. Miller

INVOICE

36881

DATE

4/14/22

PRESENTING CLINICAL SIGNS

Presented at our hospital for blood in stool, extreme lethargy, “lesions” in armpits and groin. Patient had a chewy treat last night and when owner woke up, there was blood on his blankets. Owner noticed that he was lethargic on his morning walk and had blood in his stool. Owner gave a bath and groomed him today, and noticed that he had lesions on his skin at groin and armpits (petechia). The lethargy got much worse through the day. Patient barely ate tonight and then vomited. He also had 3 bowel movements in the house, which is very unusual. Previous Health Concerns: Arthritis Current Medications: Gabapentin and a chewable arthritis med (owner unsure which)

Abnormal PE/Chem/CBC/UA Results: Eyes: OU NS, diminished to almost absent menace Cardiovascular: Grade 5/6 heart murmur Abdominal: NR, soft and non-painful Integument: Petechia all over forelimbs, abdomen, dorsum, base of tail CBC – Bas (0.15) Plt (22) CHEM – BUN (39.8) Gluc (282) ALP (569) Amy (1650) Lipase (238) EPOC – Lactate (6.31) BUN (29) Gluc (275) Radiographs – Enlarged heart; dorsal tracheal deviation; hepatomegaly 4DX – negative

ULTRASONOGRAPHIC EXAMINATION OF THE ABDOMEN

Urinary System

The **urinary bladder** was empty. Mild pseudohypertrophy noted owing to the empty bladder state and recoil effect. Foley catheter was present.

The **kidneys** presented a relatively uniform cortical hyperechogenicity when compared to the renal medulla, spleen and liver. No overt masses were noted. Corticomedullary definition was nebulous and the ratio favored the cortex slightly. The ureters were not visible and assumed to be normal. These changes are most consistent with chronic interstitial nephritis yet infiltrative disease could not be entirely ruled out without biopsy though neoplasia is not suspected. Microcystic changes noted throughout the cortices. The left kidney measured 5.66 cm. The right kidney measured 5.44 cm.

Adrenal Glands

A **left adrenal** mass was noted, measuring 3.75 cm x 2.04 cm. The **right adrenal gland** was uniform at 2.53 cm x 0.84 cm at the cranial pole and 0.67 cm at the caudal pole.

Spleen

The **spleen** presented relatively normal size and contour with multifocal hyperechoic nodular changes, most consistent with fatty deposits or lipogranulomas. These are not typically pathological. No suspicion of significant. Capsular and parenchymal integrity was normal otherwise.

Liver

The **liver** revealed increased portal markings with swollen contour. Double layered/edematous gallbladder noted.

Gastrointestinal

Examination of the **gastrointestinal tract** revealed a stomach and intestine free of stasis, of normal wall thickness, acceptable curvilinear mural detail, and peristaltic activity. Small and large intestine demonstrated normal luminal chyme and stool consistency respectively. No obstructive or overt infiltrative disease was noted. No associated abnormal lymphatic activity was noted.

Pancreas

Heterogeneous, mixed hypoechoic parenchymal **pancreatic** changes noted. Enhanced surrounding mesentery noted, consistent with pancreatitis.



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Free Abdomen

Remy Bartone

Reactive mesentery noted in the cranial abdomen.

SPECIES

Canine

- Left adrenal mass – pheochromocytoma versus adenocarcinoma
- Pancreatitis
- Cholangitis liver pattern
- Moderate degenerative renal changes with cortical cysts

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INTERPRETATION OF THE FINDINGS & FURTHER RECOMMENDATIONS

Stabilization of the pancreatitis/cholangitis presentation recommended followed by eventual left adrenalectomy. Serial blood pressure recommended. IV fluid support and treatment for the diabetic state warranted as well.

SEX

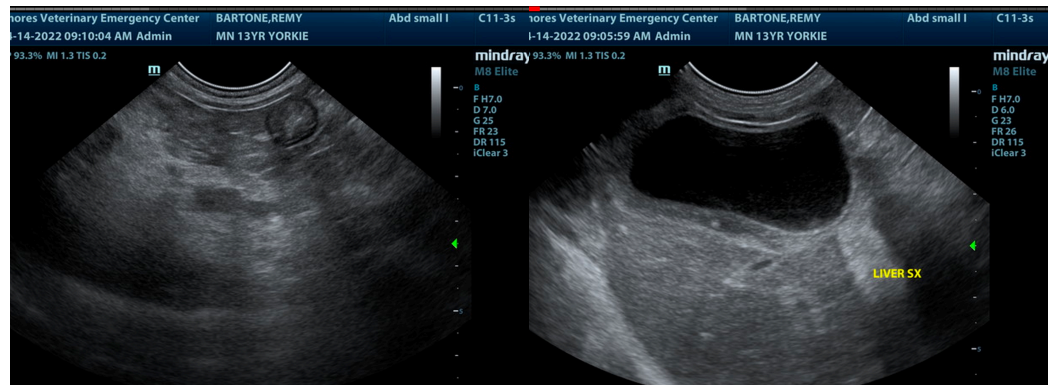
Neutered Male

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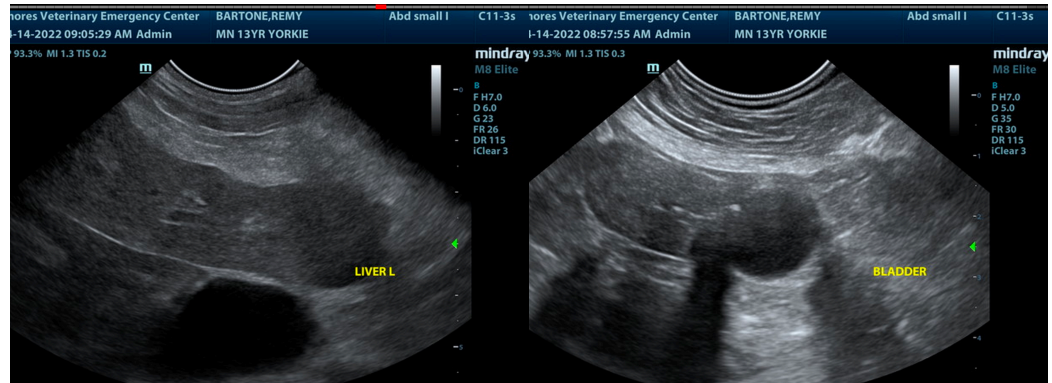
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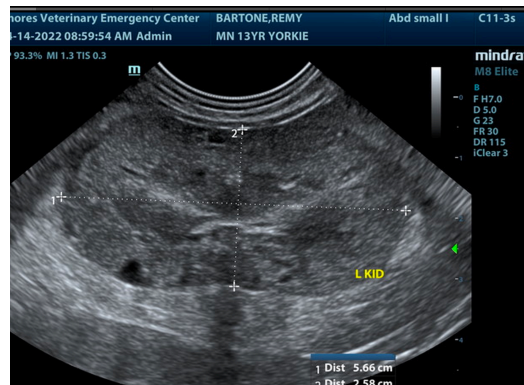
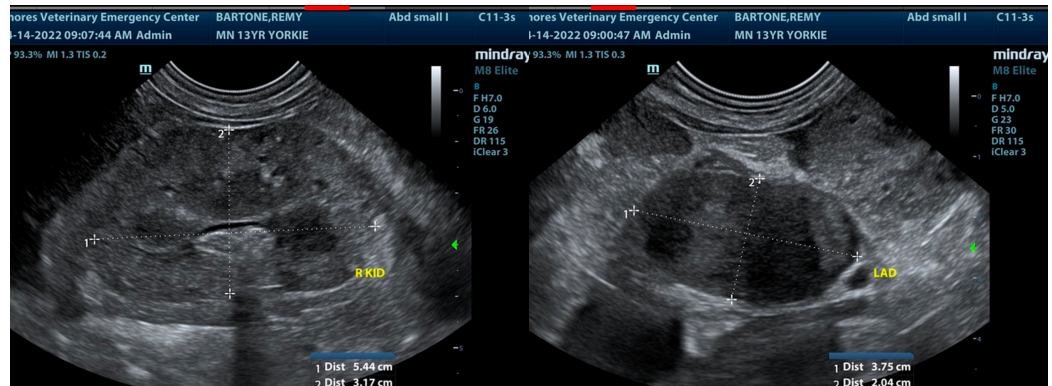
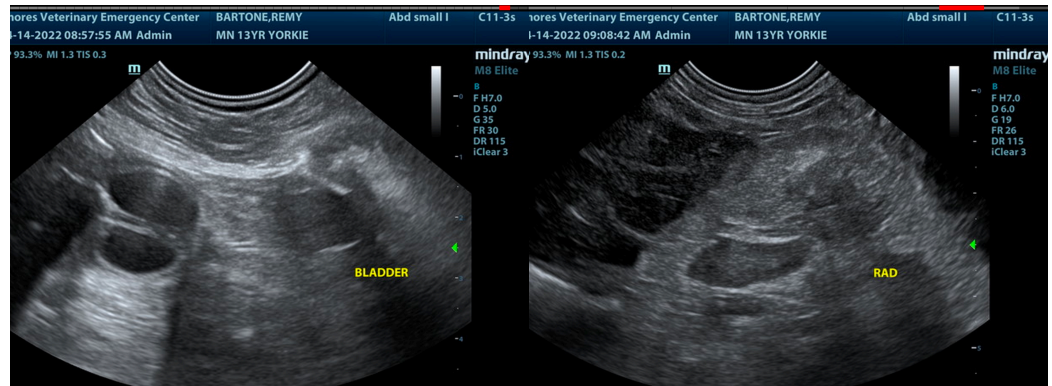
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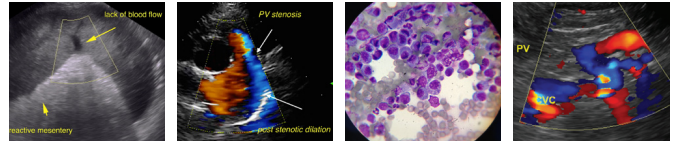


The information and recommendations provided are based on the images presented by the referring veterinarian/sonographer. No evaluation can be communicated regarding pathology that was not visible in the image/video clips provided.

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.

Eric Lindquist, DMV, DABVP, Cert. IVUSS, CEO of SonoPath.com

info@SonoPath.com



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Adrenal Tumors

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

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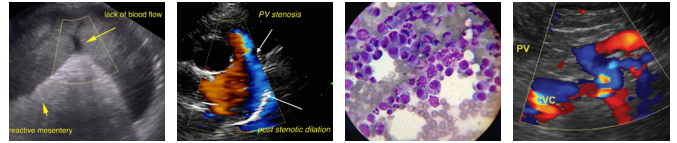
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Description: An adrenal mass is suspected when the maximum width of the adrenal gland exceeds 1.5 cm, there is loss of normal architecture or shape, or the shape or size between the affected adrenal gland and the contralateral gland is asymmetrical. The latter comprise the initial criteria for diagnosis; however, a bulbous enlargement of the cranial or caudal pole of the adrenal gland is common in dogs with no adrenal pathology and can be misinterpreted as an adrenal mass. If the suspected mass is not precipitating obvious signs (i.e., aggressive behavior), then an abdominal ultrasound should be repeated to confirm that the mass is a consistent finding before pursuing further diagnostics or surgery. Large breeds (Poodles, German Shepherds, Retrievers, and Terriers) and females appear to be overrepresented in the clinical reviews of adrenal tumors. Adrenal tumors in cats are rare with minimal information to characterize the disease. However, adrenal carcinoma and aldosterone producing tumors are the more common adrenal masses in our archived feline population. More specific information regarding this pathology may be found in the Feline Hyperaldosterone chapter.

Incidental adrenal lesions should be investigated clinically if discovered on ultrasound. Non-neoplastic adrenal lesions, such as cysts or granulomas, are very rare in dogs and cats, and the high incidence of metastatic lesions justifies a thorough hormonal screening as well as evaluation for non-adrenal neoplasms. Although incidental adrenal masses may appear to be nonfunctional at the time of diagnosis, it seems more likely that they are in fact subclinically functional. The diagnosis of functional adrenal tumors is discussed below; however, the identification of a nonfunctional, incidental adrenal mass creates a management dilemma.

Clinical Signs: Clinical signs attributable to adrenal tumors are dependent on hormone secretion type. Please see below.

Diagnostics: Cortical adrenal tumors, such as adenomas and adenocarcinomas, are responsible for 15-20% of hyperadrenocortical cases—what are commonly referred to as adrenal-dependent hyperadrenocorticism (ADH)—in dogs. The remaining tumors are the result of pituitary-dependent secretions, which give rise to pituitary-dependent hyperadrenocorticism (PDH). PDH cases tend to demonstrate bilateral hypertrophy with excessive adrenal length and, probably more importantly, width. These enlarged adrenal glands do not invade surrounding vascular structures and are defined by overstimulation resulting from excessive ACTH secretion from the pituitary gland. Yet, ADH cases are usually unilateral (bilateral in 10-20% of cases), may invade the aorta on the left or the vena cava on the right, and metastasize to the liver and lungs most frequently. Practitioners must differentiate ADH masses from hyperplastic, non-functional, benign adrenal tumors, as well as pheochromocytomas. Thus, dynamic function tests (ex. LDDS, HDDS, ACTH stimulation, ACTH baseline, urine cortisol-creatinine ratio) are essential, as is conducting routine biochemistry (ALP is elevated in more than 90% of cases) and urinalysis (true polyuria/polydipsia [PU/PD] with USG < 1.020) to determine adequately the need for surgical intervention or aggressive medical therapy. It is important to assess the following: blood pressure for hypertension; oscillating hyper- and hypotensive episodes in cases of pheochromocytomas; urine protein-creatinine ratios; and serum



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antithrombin III to determine the risk for thromboembolism. Moreover, it is essential to evaluate the entire clinical picture and objective probabilities of possessing a true hyperadrenocorticism case. This further entails ruling out other sources of PU/PD, such as primary polydipsia, renal disease, electrolyte abnormalities, infections, and diabetes insipidus or mellitus.

SPECIES

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Malignant or Benign, Functional or Non-Functional: How to Decide?

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Yorkie

In some cases, it may be difficult to determine whether the mass is malignant or benign, functional or nonfunctional, prior to surgical removal and histopathological examination. A thorough review of the clinical signs, physical examination findings, routine blood work, urine tests, and appropriate hormonal tests should be conducted to determine the functional status of an incidental adrenal mass.

SEX

Neutered Male

Malignancy is more often associated with larger masses. The larger the mass, the more likely metastasis has already occurred, in spite of a lack of detectable lesions on ultrasound and thoracic radiographs. Invasion of the mass into surrounding organs or blood vessels also supports malignancy, as does the detection of additional mass lesions with abdominal ultrasound and thoracic radiographs. Use of imaging modalities, such as CT and MRI, will likely provide additional data on the characteristics of specific adrenal lesions for use in diagnosis and treatment planning.

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Ultrasonography is the primary instrument for assessing tumor size, aggressiveness, non-capsulated versus capsulated appearance, vascular invasion, and hepatic or other metastasis. Ideally, the patient will have fasted prior to the ultrasound; one may choose to administer an enema to enhance visibility around the ascending and descending colon. Ultrasound-guided biopsy or fine needle aspiration (FNA) may be possible on the larger masses, especially on the left side; however, adjacent vascular structures often prevent the feasibility of this procedure.

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Diagnosis of the Functional Adrenal Mass:

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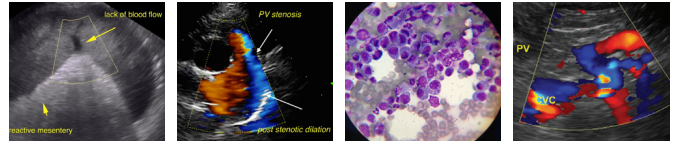
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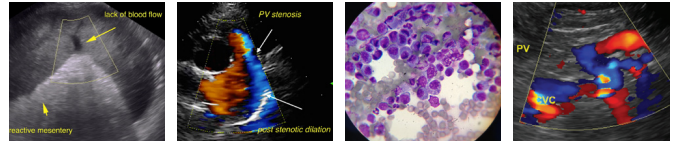
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- Cortisol-Secreting: It is very rare that a patient with hyperadrenocorticism will have a repeatable urine specific gravity greater than 1.020, so it must be determined whether the patient is truly PU/PD. If yes, then dynamic function testing is appropriate. If the patient is not truly PU/PD, then a false positive result must be considered before treatment is initiated, as the resulting hypoadrenocorticism can be life threatening. Other causes of dysuria, such as occult urinary tract infection, must then be considered. The most common functional adrenal tumor identified in dogs and cats results in hyperadrenocorticism. Approximately 15% of hyperadrenocorticism cases will be caused by a functional adrenal tumor, of which 50% of these will be malignant.
 - Clinical signs can include: PU/PD; polyphagia; abdominal distention; bilaterally symmetrical truncal alopecia; delayed fur regrowth; hyperpigmentation; comedones; calcinosis cutis; excessive bruising; poor wound healing; ectopic calcification of kidneys and blood vessel walls; pyodermas; muscle weakness; exercise intolerance; hypertension; and panting.
 - Ultrasound usually reveals a small or atrophied contralateral adrenal gland as a result of suppressed pituitary ACTH secretion. Ten to twenty percent of cases have bilateral disease. Adenomas of the adrenal gland are generally less than 2 cm in diameter, and carcinomas can be any size (often they are > 2 cm). Calcification does not appear to be predictive for either adenoma or carcinoma.
 - Specific biochemical tests: Urine cortisol-creatinine ratio, ACTH stimulation test, and LDDS test.
- Catecholamine-Producing: Pheochromocytoma is a tumor derived from the chromaffin cells of the adrenal medulla; it is relatively common in dogs, but quite rare in cats. These cases should be considered malignant until proven otherwise. Invasion/entrapment/compression of the caudal vena cava is common. Mural invasion or luminal narrowing of the aorta, renal vessels, adrenal vessels, and hepatic veins may also occur.
 - Clinical signs associated with this type of tumor are usually related to the invasion of local structures, metastases, or the secretion of catecholamines. The most common clinical signs of excess catecholamines



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SPECIES	<ul style="list-style-type: none"> ○ Ultrasound: The contralateral adrenal gland is usually normal in size and shape. Pheochromocytomas do not typically calcify. ○ Tests: Many of the clinical signs and blood pressure alterations are similar for pheochromocytoma and ADH. It is therefore important to rule out ADH before focusing on pheochromocytoma. The diagnosis prior to surgery is primarily one of exclusion. Specific hormonal tests, such as those that measure urinary catecholamine concentrations or their metabolites, are not routinely performed.
Canine	
BREED	<ul style="list-style-type: none"> ● Aldosterone-Secreting (rare in dogs and cats): <ul style="list-style-type: none"> ○ Clinical signs (Conn's Syndrome) are related to excessive secretion of aldosterone, which causes sodium retention and potassium depletion. The resulting symptoms include lethargy, weakness, mild hypernatremia, severe hypokalemia (usually < 3.0 mEq/L), and systemic hypertension. ○ Ultrasound usually reveals a normal contralateral adrenal gland. ○ Tests: Documenting increased plasma aldosterone concentrations before and after ACTH administration is a means of confirming the diagnosis. If weakness and severe hypokalemia are present, plasma aldosterone concentrations can be measured along with plasma cortisol concentrations during the ACTH stimulation test. ● Progesterone-Secreting: Although a functional tumor arising from the zona reticularis of the adrenal cortex could secrete excessive amounts of estrogen, progesterone, or testosterone, to date only progesterone-secreting adrenocortical tumors in cats have been documented. <ul style="list-style-type: none"> ○ Clinical signs include: diabetes mellitus and feline fragile skin syndrome, which is characterized by progressively worsening dermal and epidermal atrophy, patchy endocrine alopecia, and easily torn skin. ○ Ultrasound usually reveals a normal contralateral adrenal gland. ○ Tests: Diagnosis requires documenting an increased plasma progesterone concentration. The clinical features mimic feline hyperadrenocorticism, which is the primary differential diagnosis. Pituitary-adrenocortical axis test results are normal to suppressed in cats with progesterone-secreting adrenal tumors. ● Deoxycorticosterone-Secreting (rare): <ul style="list-style-type: none"> ○ Clinical signs are related to mineralocorticoid activity and include weakness, marked hypokalemia, and systemic hypertension. ○ Tests: Increased plasma deoxycorticosterone and non-detectable plasma aldosterone concentrations have been documented in dogs. ● 17-OH-progesterone-Secreting (rare): <ul style="list-style-type: none"> ○ Clinical signs are similar to hyperadrenocorticism. ○ Tests: Pre- and post-ACTH stimulation plasma 17-OH-progesterone concentrations will be increased.
Yorkie	
SEX	
Neutered Male	
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REFERRING VET	<p>When a cortisol-producing adrenal tumor has been documented, medical therapy with trilostane (5-20mg/kg PO Q24hr) or mitotane (25-50 mg/kg PO Q24hr for 10 days, then every 4-7 days) should be considered.</p> <p>The biggest dilemma is whether to perform an adrenalectomy if hormonal tests for hyperadrenocorticism and serum electrolyte concentrations are normal, and clinical signs and systemic hypertension suggestive of pheochromocytoma are not present.</p>
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INVOICE	<p>An aggressive approach—adrenalectomy—is based on the assumption that the mass is malignant until proven otherwise and should be removed before metastasis has occurred. In theory, this approach would offer the best chance for long-term survival; however, the age of the patient, the size of the mass, the presence of concurrent diseases, the level of invasion into other organs, and the probability that metastases already exist should factor into the decision. Poor surgical candidates generally include: dogs compromised from the effects of hypercortisolis; older animals; animals with concurrent disease; those for whom invasion has been aggressive and surgical or post-surgical complications are likely; animals with very large masses that have likely already metastasized; and those with documented potential metastatic disease. In addition, adrenalectomy may not be indicated when the</p>
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mass is small (< 3 cm diameter) and nonfunctional, and the patient is healthy. Reports suggest that there is an approximate 45% success rate of surgical resection of adrenal masses, with a positive prognosis inversely proportionate to tumor size.

SPECIES

Canine

In cases of concurrent hepatic nodular changes, liver biopsy samples can be obtained at surgery in cases of suspicious lesions visualized by ultrasound. Hyperadrenocorticism often causes benign nodular hyperplasia of the liver and should not be automatically interpreted as a sign of hepatic metastasis during ultrasonographic examination. Rather, suspect lesions should be confirmed and biopsied either at surgery or via ultrasound-guided FNA or core biopsy. Post-operative complications include delayed wound healing due to excessive corticoid circulation and wasting, hemorrhage, sepsis, and thromboembolism.

BREED

Yorkie

When surgery is a risk and a functional adrenal tumor has been documented, medical therapy, as outlined above, should be considered. Medical therapy will not impede metastatic events. An alternative approach in these cases is to determine the rate of growth of the mass by repeating abdominal ultrasounds initially at 2, 4, and 6 months. If the adrenal mass does not change in size, the time between ultrasound evaluations can be increased to every 4-6 months; however, if the adrenal mass is increasing in size, adrenalectomy should be considered.

SEX

Neutered Male

Conclusion: Any incidentally discovered adrenal tumor warrants investigation into functionality and metastasis. The course of treatment for each case depends largely on which hormones are secreted by the adrenal tumor. Each case should be carefully evaluated on an individual basis before adrenalectomy is considered for aggressive tumors.

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