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INTRODUCTION

The anatomy of the kidney is conserved among mammals. The kidneys are located within the sublumbar peritoneal space, with the right kidney positioned cranially to the left. In female rodents, there is a separate external orifice for the urinary tract and the reproductive tract. In male and female sugar gliders (Petaurus breviceps), the urinary tract terminates into a cloaca. Male chinchillas (Chinchilla lanigera), guinea pigs (Cavia porcellus), rats (Rattus norvegicus), mice (Mus musculus), hamsters, gerbils, and ferrets (Mustela putorius furo) have an os penis or baculum.

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There are a few histologic differences of the kidney, including the number of papilla and length of the nephrons. Most differences are related to the natural environment of the animal. These differences are noted between desert-dwelling species requiring optimal ability to concentrate urine, and other species in which water deprivation is not common. Of clinical significance is the specific gravity of urine in desert-dwelling species, which is generally higher.\(^2\)

**FERRET**

**Common Problems**

The most common problems affecting the ferret urinary system described in the literature are Aleutian disease of the kidney, renal tumors, renal cysts, urolithiasis, and bacterial cystitis. In one author’s database (DRR), hydronephrosis, nephritis (all causes), and renal mineralization are the most common lesions of submitted kidneys.

**Infectious/inflammatory**

Aleutian disease is caused by a parvovirus.\(^3\) The lesions are immune mediated and vary in severity. Gross lesions may be minimal and nonspecific. Emaciation and organ enlargement can be seen. In the kidney, membranous glomerulonephritis is common and a lymphocytic-plasmacytic inflammatory infiltrate is seen in the interstitium. This infection is uncommon in ferrets and the mortality is generally low.

Systemic coronavirus can also affect the kidneys, and the primary lesion is granulomatous inflammation.\(^4\) In a study showing ultrasonography results in ferrets with systemic coronavirus, nephromegaly was found in 4 out of 11 patients.\(^5\)

Canine distemper is essentially 100% fatal. Affected ferrets present with neurologic clinical signs, bronchopneumonia, hyperkeratosis of the planum nasale and footpads, and a papular rash starting on the chin. However, the most productive tissues to evaluate for viral inclusions are the urinary bladder, renal pelvis, and biliary epithelium, and in suspected cases these tissues should be submitted (Fig. 1). Immunohistochemistry can be applied for confirmation.

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*Fig. 1.* Ferret (*M. putorius furo*) urinary bladder with intracytoplasmic inclusions of a canine distemper infection. Confirmed with immunohistochemistry. Black arrows indicate a few of the intracytoplasmic eosinophilic viral inclusions (HE stain, 20× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).
Bacterial nephritis and pyelonephritis occur in ferrets and can be severe. These conditions are thought to be caused by ascending infections of the lower urinary tract. A variety of organisms can be the cause. Grossly, the kidney may be abscessed and histologically there is necrosis and a neutrophilic infiltrate. Organisms may be seen but culture is necessary for a definitive diagnosis. Bacterial infection of the kidney, prostate, or urinary bladder can result in inflammatory cells (often undergoing degeneration) and bacteria in the urine.

**Degenerative/congenital**

Nephrosis is considered a degenerative disease of geriatric ferrets, and is often an incidental finding. Nephrosis is characterized by damage of tubular epithelial cells. It is a nonspecific lesion that can have many causes, including ischemia, toxic exposure, and shock.

Nephrocalcinosis is reported more frequently in European pet ferrets than laboratory ferrets, and lesions include calcium deposition in the renal tubules. The cause of renal mineralization is usually not determined. Soft tissue mineralization is usually considered dystrophic (secondary to tissue injury) or metastatic (associated with excessive calcium in the blood), but neither cause is obvious in most cases seen at necropsy. The lesion is usually in the renal papilla. Rare cases of lymphoma-associated hypercalcemia have also resulted in renal dystrophic mineralization. The mechanism is most likely azotemia and the nephrotoxic effects of prolonged hypercalcemia.

Renal cysts are a common and usually incidental finding in the ferret (Fig. 2). A 17-year retrospective analysis of 54 ferrets with cystic kidney disease noted that 69% had renal cysts, and 26% primary polycystic disease. Cysts may be single or multiple and may be present in the cortex of 1 or both kidneys. It is speculated that acquired cysts arise from gradual distention of the nephron, caused by obstruction by exudates or fibrosis tissue, although the cause is seldom determined. Rare cases of true polycystic disease have been reported in the ferret. This congenital lesion may result in markedly enlarged cystic kidneys that fill the posterior abdomen. Many small cysts throughout the cortex and medulla characterize polycystic kidneys.

![Fig. 2. Renal cysts within the cortex of the ferret (M putorius furo) kidney (HE stain, 1× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).](image-url)
Another rare congenital lesion is an extramural ectopic ureter. This condition is the most common congenital cause of urinary incontinence in domestic canines. Young ferrets present with urinary incontinence and urine scalding around perineal and inguinal areas. The ectopic ureter can be visualized via excretory urography. In 1 case, nephroureterectomy was chosen instead of ureteral transplant. The ureter progressed extramurally to enter the distal urethra, caudal to the urethral sphincter, or vagina (on contrast).

**Neoplastic**

The most common renal neoplasm is malignant lymphoma. Lymphoma usually involves multiple organs and can be severe in the kidney. Grossly, there is variable replacement and distortion of the kidney. Histology is typical, with a diffuse monomorphic sheet of lymphoid cells. Impression smears can often provide a diagnosis. Other neoplasms have been reported, although primary tumors of the urinary tract system are uncommon. From 1 study of 574 ferret tumors, only 6 urinary tumors were recognized. There were 3 renal carcinomas and 3 unspecified renal tumors. Another study evaluating 856 ferret tumors identified only 1 renal tumor listed as a sarcoma, not otherwise specified, and lymphoma involving the kidney. One renal adenocarcinoma is described in detail collected from an adult neutered female. This pleomorphic renal adenocarcinoma had multiple metastases to the lung, liver, greater omentum, right renal pelvis, and systemic lymph nodes. The tumor cells were pleomorphic with a large number of giant cells and arranged in tubular and cystic patterns. It was confirmed of renal origin by immunohistochemistry with strong positive CD10 and cytokeratin. Two cases of transitional cell carcinoma arising from the renal pelvis, with 1 completely replacing the kidney, have been reported. The presenting complaint was hematuria.

**Obstructive disorders**

**Urolithiasis** The clinical signs of urolithiasis include stranguria and dysuria. Uroliths in ferrets commonly consist of sterile struvite or magnesium ammonium phosphate. Based on 1 study evaluating 272 cases, neutered males have a significantly increased risk of developing sterile struvite urolithiasis. These uroliths are more likely to be retrieved from the lower urinary tract than from the upper urinary tract. Stone formation is suspected to be caused by a vegetable-based protein diet that increases urine pH, leading to struvite stone formation.

Cystine uroliths are uncommon, with a reported incidence of 16% in 1 study evaluating 70 cases. Again, males are predisposed to urolithiasis, possibly because the distal portion of the urethra has a J-shaped bend with a decreased diameter of the urethra. All cystine uroliths have been found in the lower urinary tract. The uroliths are ovoid and smooth, light yellow to tan, and range in number from 1 to more than 100. Cystine uroliths are composed of nonessential sulfur-containing amino acid; there may be a familial cause based on studies in other mammals.

Calcium oxalates are the next most common type of urolith.

**Prostatic disease** Other causes of urinary obstruction in the male ferret are abnormalities of the prostate caused by adrenocortical disease. Males can develop prostatic hyperplasia, cysts, and abscesses. Prostatic enlargement is caused by increased levels of the hormones estradiol, testosterone, and 17-hydroxyprogesterone produced by the neoplastic adrenal cortex from an adrenal gland tumor. Some animals develop concurrent urinary tract infection, including small urethral or cystic calculi. Urethral blockages can also be caused by sloughed proteinaceous debris from
squamous metaplasia of prostatic tissue and by small struvite calculi, which develop from the increase in pH caused by bacterial cystitis.²⁰

Preputial tumors in male ferrets can also result in urethral obstruction. These aggressive tumors are poorly responsive to both surgery and radiation therapy. Nevertheless, wide surgical resection has been recommended, although the procedure may necessitate partial or total penile resection (Fig. 3).²¹

**Treatment**

Any disease of the kidneys can result in renal insufficiency or failure. Symptoms may include polyuria/polydipsia (PU/PD), weight loss, weakness, nausea, and decreased appetite, as can be seen in other traditional pet species. Physical examination findings are also consistent and often include dehydration, although hydration status can be challenging to determine by testing skin turgor in patients experiencing rapid weight loss. Some clinicians have anecdotally reported halitosis and oral ulcers in association with renal failure, but this is poorly documented.

Treatment protocols are based on those for traditional pet species, and include fluid diuresis for renal insufficiency and failure, antibiotic therapy for bacterial nephritis, and basic surgical approaches to the kidney.

Treatment of urinary tract disease follows guidelines established for traditional pet species, such as treatment of bacterial disease based on results of culture and sensitivity. General principles for treatment of renal insufficiency and renal failure are described later in this article. Treatment of adrenocortical disease is well described in any current literature about ferret endocrine diseases. Large associated prostatic cysts have been managed with marsupialization or omentalization of the cyst.²⁰

Successful nephrotomy for removal of unilateral renoliths has been reported in the ferret.²²

**RABBIT**

**Common Problems**

The most common causes of renal disease reported in the literature are *Encephalitozoon cuniculi*, chronic renal failure in older rabbits (*Oryctolagus cuniculus*), and urolithiasis. Other renal conditions seen in rabbits include bacterial nephritis, hydronephrosis, and papillary necrosis with renal mineralization.

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Fig. 3. Preputial tumor on a ferret (*M putorius furo*), prepped for surgical removal. (Courtesy of S. A. Kelleher, DVM, Deerfield Beach, FL).
Infectious/inflammatory
The most common renal disease of rabbits is infection caused by *E cuniculi*.\(^{23–25}\) Although disease is often occult, rabbits with severe *E cuniculi*–associated renal disease may present with PU/PD and a host of other nonspecific signs, such as decreased appetite and weight loss, dehydration, and lethargy. In some rabbits, only renal changes are seen at necropsy. Grossly, there may be no lesions or the kidneys may be pitted. On histology, there is a chronic active lesion, which may be granulomatous or pyogranulomatous. Aggregates of organisms are found rarely in or around tubules as small, basophilic structures (Fig. 4).

Rare cases of polypoid cystitis and urethritis have been described in rabbits. The masses are white to transparent, spheroidal masses protruding into the lumen. On histology, the polyps are lined by variably hyperplastic to attenuated transitional epithelium over cores of edematous inflamed fibrovascular stroma. There may be transitional epithelium lining cysts within the stroma. These polyps are suspected to arise from a combination of inflammation and a hyperplastic reaction to chronic irritation of the urinary mucosa. Bacterial isolates have included *Proteus* and *Enterococcus* spp. The clinical signs associated with polypoid cystitis include perineal scalding, urinary sludge, and possibly hematuria.\(^{26}\) Hematuria is frequently described as a clinical sign with obstructive urolithiasis, cystitis, bladder polyps, and pyelonephritis.\(^{27}\) Red urine in rabbits may not necessarily be hematuria but may be the more common, and normal, porphyrin pigments. Urine dipstick is an easy method to differentiate porphyrinuria from hematuria.

Degenerative/congenital
Chronic renal disease or end-stage kidney disease seem to be common in older rabbits. Rabbits with renal failure can have lesions similar to those seen in old rats. The

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**Fig. 4.** *E cuniculi* microbes are within the tubular epithelial cells and these are sloughing into the lumen of the tubules. The black arrows indicate intratubular and intraepithelial densely packed microbes, which are small round/oval amphophilic structures. Samples from a dwarf rabbit (*O cuniculus*) (HE stain, 40× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).
kidneys are grossly scarred and histologically there is marked interstitial fibrosis and glomerulosclerosis. This condition may be a sequela of encephalitozoon infection, but the cause is usually not determined.

Renal cysts are uncommon in rabbits. In a few reports, some cases were speculated to be inherited and to resemble human polycystic kidney disease. In experimental studies, methylprednisolone acetate has been injected into newborn rabbits to induce polycystic kidneys. The few cases of renal cysts (7 out of 796 rabbit renal submissions) in 1 author’s database (DRR) have been in older rabbits (>5 years) and associated with lesions of chronic renal disease.

**Neoplasia**

Tumors of the urinary tract are rare in rabbits, with most involving the kidney. These tumors include benign embryonal nephroma, renal carcinoma, renal adenocarcinoma, nephroblastoma, and kidney hamartoma. A case of renal and ureteral transitional cell carcinoma produced hydronephrosis. A urinary bladder transitional cell carcinoma in a female rabbit presented with acute hematuria, partial vaginal prolapse, and inappropriate urination. The gross appearance was a nodular cystic mass at the cranial central apex of the urinary bladder.

**Obstructive disorders**

**Urolithiasis and urinary sludge** Uroliths are described in the kidney and urinary bladder of rabbits. Renoliths can be mild with few stones, or severe and produce renal megaly and complete destruction the kidney. Uroliths and urinary sludge in rabbits are usually composed of various calcium salts, predominately calcium carbonate. From a study evaluating more than 100 rabbit uroliths, the most common were calcium carbonate (69.4%), compound (23%), and mixed (3.3%). Radiographs are usually confirmatory because calcium carbonate is radiopaque. The predominance of calcium-based uroliths is most likely caused by the unique calcium metabolism of rabbits. Excess dietary calcium is absorbed unregulated, resulting in increased plasma calcium concentrations and excretion of excess calcium in the urine. Therefore, serum and urinary calcium levels are related to dietary intake. Urinary calcium can bind with other constituents when conditions are met for crystallization, such as dehydration, changes in pH, and urine

![Fig. 5. Rabbit (O cuniculus) nephroblastoma is effacing the kidney (HE stain, 10× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).](image-url)
retention, to form both the sludge (which can appear as thick white to yellow material) and uroliths. Abnormal accumulations of calcium (sludge) are more common in older rabbits, possibly because of decreased mobility or water consumption.\textsuperscript{34} There are rare reports of uroliths composed of silica, struvite, and calcium sulfate dehydrate or gypsum.\textsuperscript{35} The gypsum urolith was associated with the rabbit eating gypsum-based plaster that was covered by a white paint containing barium sulfate.

**Hernia**

Hernias can be classified by their anatomic location (body cavities or external, such as inguinal or femoral), cause (true hernia [through an anatomic structure and surrounded by peritoneum] or false hernia [traumatic through induced defects]), and contents.

Inguinal or scrotal hernias develop when abdominal contents, such as the urinary bladder, enter the inguinal canal or scrotum through a wide or defective inguinal canal. In male rabbits, the inguinal rings are open throughout life, increasing the risk of herniation. The typical appearance of bladder herniation is a fluid-filled mass in the scrotum and no bladder palpable in the abdomen. Some cases have concurrent

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**Fig. 6.** (A) Bilateral nephrolithiasis in a rabbit (*O cuniculus*). The radiographs highlight the radiodense materials within the pelvis of the both kidneys. (B) Lateral view. (Courtesy of A. Lennox, DVM, DABVP (Avian Practice, Exotic Companion Mammal Practice), Diplomate ECZM (Small Mammal), Indianapolis, IN).

**Fig. 7.** Uroliths in the bladder can be of variable size and shape; although many are smooth, this urolith contained sharp points. The rabbit (*O cuniculus*) presented with anorexia and hunched posture, suggesting discomfort. (Courtesy of A. Lennox, DVM, DABVP (Avian Practice, Exotic Companion Mammal Practice), Diplomate ECZM (Small Mammal), Indianapolis, IN).
lesions of urinary sediment and multiple uroliths; 1 case featured a testicular tumor on the opposite side. Inguinal herniation of the bladder seems uncommon in female rabbits (Fig. 8). Trauma was suspected to be the cause in 1 case of a female rabbit presenting with hematuria.

Female rabbits may also evert the urinary bladder through the urethra. Recent kindling or a history of multiple litters is common. Rabbits present with a vaginal mass, which is the mucosal surface of the everted bladder. Rarely, males may evert and prolapse the urinary bladder. This finding seems more likely in castrated males because penis length is reduced after castration.

Another unusual presentation is diaphragmatic herniation of retroperitoneal fat and a kidney. A case of kidney herniation occurred in conjunction with a diaphragmatic lipoma.

**Treatment**

General management of renal insufficiency and renal failure is described later.

**Encephalitozoonosis** Treatment of *E cuniculi* is problematic because there is a lack of well-controlled scientific studies linking various treatment protocols with documented clearing of organisms. The problem relates to difficulties in establishing a diagnosis, and the observation that many affected rabbits recover spontaneously without treatment. Rabbits with severe renal disease secondary to *E cuniculi* may not harbor renal organisms, and treatment is focused on managing acute or chronic renal failure. Treatment of *E cuniculi* has relied on benzimidazoles, such as fenbendazole, but recently severe disease and deaths have been reported in rabbits treated with this class of drugs. Benzimidazoles are radiomimetic and toxicity has been observed in many species; Graham and colleagues recently reported deaths in 13 rabbits treated with various dosages of several drugs, including fenbendazole, albendazole, and oxibendazole. Clinical signs in rabbits were nonspecific and histopathology showed depletion of all bone marrow cell lines. For this reason, treatment should not be approached lightly, and serial monitoring of the complete blood count may be beneficial.

**Surgical management of renal disease** Various surgical approaches to the kidney have been described. Martorell and colleagues described a lateral flank approach

Fig. 8. Inguinal herniation of the bladder in a female rabbit (*O cuniculus*). The bladder is filled with microuroliths (bladder sludge), which are acting as natural contrast. Surgery to replace the bladder and correct the herniation was successful. (A) Lateral and (B) frontal view. (Courtesy of A. Lennox, DVM, DABVP (Avian Practice, Exotic Companion Mammal Practice), Diplomate ECZM (Small Mammal), Indianapolis, IN).
for access to the kidney for removal of uroliths; 1 author (AML) also found this approach simple and straightforward compared with a traditional midline abdominal approach (Fig. 9). Nephrectomy, performed through a ventral abdominal approach, was reported to manage hydronephrosis in a pet rabbit and to remove a renal carcinoma. Cases of diaphragmatic herniation were successfully treated by replacing the herniated kidney and fat followed by diaphragmatic herniorrhaphy, and use of polypropylene mesh.

Management of bladder sludge Strategies for managing urinary sludge include encouraging water consumption and exercise paired with dietary modification as needed. Accumulation of sludge can be temporarily managed by bladder catheterization and flushing, or sometimes by gently agitating the bladder to manually resuspend the mineral, followed by repeated expression of urine.

GUINEA PIG

Common Problems

Urolithiasis is commonly reported in the guinea pig. This condition usually involves the lower urinary tract. Inflammatory, degenerative, and other conditions are also reported.

Infectious/inflammatory

Chronic nephritis, interstitial nephritis, and pyelonephritis have been reported. In these conditions the kidneys may be irregular grossly and the ureters may be dilated in pyelonephritis. On histology, the inflammation depends on the cause and the duration of the disease.

Cystitis in guinea pigs is common. Guinea pigs, like chinchillas and degus, have adapted to a semiarid habitat with varying, species-specific adaptations to water deprivation. However, several diseases have been linked to insufficient water intake, such as cystitis, urolithiasis, or obstipation. A decrease in urination frequency may favor bacterial infection and formation of uroliths. The clinical signs of bacterial cystitis include dysuria, polyuria, hematuria, lethargy, or urethral discharge. The urinary bladder may be painful on palpation, small, firm, with a thickened wall. The most common isolate in a review series of adult female guinea pigs was Escherichia coli. Relapses were common. Two guinea pigs were diagnosed with encrusted cystitis (further description in relation to hedgehogs later).

Fig. 9. Unilateral flank approach nephrotomy allowed removal of uroliths from the right kidney. (Courtesy of A. Lennox, DVM, DABVP (Avian Practice, Exotic Companion Mammal Practice), Diplomate ECZM (Small Mammal), Indianapolis, IN).
Degenerative/congenital
Renal cysts are a rarely reported and usually incidental finding. Given the few reports of renal cysts in guinea pigs, it seems most cases are acquired cysts associated with chronic renal disease. One author (DRR) has seen 21 cases in adult guinea pigs (3–7 years of age), all with chronic renal lesions (fibrosing interstitial nephritis) out of 430 case submissions with kidneys (4.8%).

Chronic (segmental) nephrosclerosis is renal scarring of undetermined cause, which may be the result of vascular disease, infection, or immune-mediated disease. Grossly, the renal cortex is irregularly pitted and histologically there is interstitial fibrosis, variable glomerulosclerosis, tubular dilatation, and variable mononuclear inflammation (Fig. 10).

Renal mineralization can be incidental or associated with renal failure, resulting in uremia and soft tissue mineralization. The mechanism is complex but involves tissue death and secondary mineralization. In guinea pigs, exposure to excessive vitamin D is a common mechanism if renal failure is not present. Such exposure occurs when feeding guinea pigs a commercial diet with errors in formulation or commercial rodent diets. Commercial rodent diets are formulated with more vitamin D than is safe for guinea pigs.

Neoplasia
Primary tumors are rare. Lymphoma occasionally involves the kidneys. Grossly, there is a diffuse white-gray infiltrate that replaces normal parenchyma. On histology, this is composed of immature lymphoid cells.

Fig. 10. Gross view of chronic renal disease in a guinea pig (C porcellus). Bands of blotchy white areas radiate out from the pelvis. The histology findings were of severe diffuse fibrosing lymphoplasmacytic interstitial nephritis with periglomerular sclerosis. (Courtesy of S. Brenner, MS, DVM, San Mateo, CA).
**Obstructive disorders**

Uroliths may be present from the renal pelvis to the urinary bladder. If in the ureter or bladder, there may be ureteral dilatation and renal swelling or shrinkage. The most common urolith is calcium carbonate, which is radiopaque. Similar to rabbits and other hindgut-fermenting herbivores, guinea pigs excrete excess calcium and they can eliminate up to 75% of their absorbed calcium via urine. Urine specific gravity and pH highly influence crystallization. Herbivore urine is commonly alkaline, which complicates dietary modification efforts to change urine pH and reduce urolith formation.

**Treatment**

Strategies for treatment of acute and chronic renal failure are described later. Surgery of the kidney has not been described in pet guinea pigs. The urethra of the female guinea pig is short and wide. Uroliths in the urethra and in the vaginal vestibulum can be visualized and removed manually with the aid of the Lone Star retractor.

**HAMSTER**

**Common Problems**

Primary renal disease is uncommonly reported in pet hamsters, and mostly includes infectious and degenerative causes.

**Infectious/inflammatory**

Sporadic cases of bacterial cystitis and pyelonephritis are seen. Gross changes include thickening and exudation in the bladder and yellow-white foci of necrosis and inflammation in the kidney. Culture is usually necessary to establish a definitive cause.

**Degenerative/congenital**

The most common problem seen in the kidneys is amyloidosis, which has been reported in more than 50% of hamster necropsies in some colonies. Amyloid is an insoluble pathologic proteinaceous substance, deposited between cells in various tissues and organs of the body. Systemic amyloidosis can be classified as primary (amyloid light chain), secondary (amyloid associated), or familial. Secondary amyloidosis is most commonly described in animals and has been described as a reaction to diverse inflammatory stimuli. Amyloid deposits are more common in female hamsters more than 18 months of age. Deposits can also be seen in almost any other organ. Amyloidosis can be associated with nephritic syndrome. Grossly, the kidneys are pale and irregular. Microscopically, amyloid is eosinophilic to amphophilic and can be seen in glomeruli, interstitium, or both.

Glomerulonephropathy occurs in hamsters as well as other rodents. Progressive nephropathy can result in renal failure. Grossly, kidneys are pitted and variably shrunken, and histologically there may be variable stages of mesangial and glomerular capillary thickening, periglomerular sclerosis, and variable interstitial nephritis. This condition has been described in both Chinese (Cricetulus griseus) and Siberian hamsters (Phodopus sungorus).

**Neoplasia**

Primary tumors are rare in hamsters. Two cases of renal adenoma have been reported in aged Siberian hamsters. One author (DRR) has seen 2 cases of lymphoma involving the kidney as well as 1 case of a renal adenoma out of 60 submissions that included the kidney.
**Obstructive disorders**

Uroliths are also uncommon in hamsters. From 1 study evaluating only 14 samples, the most common uroliths were calcium phosphate (28.6%), compound (28.6%), and calcium oxalate (21.4%).

**Treatment**

Strategies for treatment of acute and chronic renal failure are described later. Surgery of the kidney has not been described in pet hamsters.

**MOUSE**

**Common Problems**

Renal disease is common in mice. Some conditions are strain related, and determining the genetic heritage of a pet mouse (if possible) may be of value in diagnosis. Amyloidosis is seen primarily in older animals.

**Degenerative/congenital**

Chronic progressive nephropathy or chronic nephritis in mice is similar to the condition in rats and other rodents. Pathogenesis is unknown. Interstitial nephritis and pyelonephritis are also reported.

Grossly, kidneys affected by these conditions have an irregular pitted cortex. Microscopically, there is glomerular sclerosis, tubular dilatation and degeneration, interstitial inflammation, and fibrosis. Lesions vary in severity according to their duration. In cases of klossiellisis, organisms can usually be seen on histology and may not be associated with significant inflammation. This organism is a parasitic protozoa of the phylum Apicomplexa. Species in this genus infect the renal tract of mammals and intestinal tract of snakes. This genus of the phylum Apicomplexa is unusual in having only a single host in its life cycle.

Polycystic kidneys occur in BALB/c mice, which are used as a model for human disease. Cysts vary in size and number, but the condition is progressive and results in loss of renal function and early mortality. The renal surface is irregular and fluid-filled cysts are visible. Cysts are also obvious in cut sections. Dilated tubules are seen histologically.

**Neoplasia**

Renal neoplasia occurs sporadically. Malignant lymphoma can efface the kidney and carcinomas are occasionally seen. Gross lesions are not specific. On histology, lymphoma is typical and carcinomas are usually composed of poorly differentiated cells with a loss of organized structure.

**Obstructive disorders**

Hydronephrosis can be unilateral or bilateral. Unilateral hydronephrosis is often an incidental necropsy finding and may be strain related or secondary to obstruction or inflammation of the renal pelvis. It is usually progressive and grossly there is variable dilatation of the renal pelvis with loss of renal parenchyma. One cause of obstructive hydronephrosis in male mice, as well as in rats and guinea pigs, is the retrograde movement of seminal plugs in the urethra (Fig. 11). These seminal vesicle secretions may coagulate and become impacted in the urethra and urinary bladder.

**RAT**

**Common Problems**

Common renal conditions reported include hydronephrosis, polycystic kidneys, and pyelonephritis, as well as neoplasms.
Infectious/inflammatory

*Trichosomoides crassicauda* is a nematode that affects the urinary tract of rats. Adult worms can be seen in the renal pelvis and urinary bladder, but the condition is usually asymptomatic. Eggs are passed in the urine. Nematodes are present in the epithelium and there is variable inflammation (Fig. 12).72

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**Fig. 11.** Seminal plugs within the lumen of the urethra in a male guinea pig (*C. porcellus*). The acellular coagulum is within the lumen. (HE stain, 20× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).

**Infectious/inflammatory**

*Trichosomoides crassicauda* is a nematode that affects the urinary tract of rats. Adult worms can be seen in the renal pelvis and urinary bladder, but the condition is usually asymptomatic. Eggs are passed in the urine. Nematodes are present in the epithelium and there is variable inflammation (Fig. 12).72

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**Fig. 12.** *T. crassicauda*, the rat (*R. norvegicus*) urinary bladder nematode can be found in the lumen as well as in the mucosal epithelium (black arrows). (HE stain, 10× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).
Degenerative/congenital
The most common renal disease in rats is chronic progressive nephropathy or old rat nephropathy.73–75 The incidence can be up to 75% in affected strains. This disease affects older animals and is more common and severe in males, with a much higher incidence in S-D and Fischer 344 rats. High-protein diets are a factor in pathogenesis, as are immunologic factors, such as mesangial deposition of immunoglobulin M, and high levels of prolactin. Lesions of progressive nephropathy are characterized by irregular and pitted renal cortices. The cut surface is also irregular. Microscopic changes vary with duration. Glomeruli are thickened, tubules are dilated and may contain proteinaceous fluid, and there is interstitial fibrosis and inflammation.

Nephrocalcinosis is seen, usually as an incidental finding but also with diets low in magnesium and/or high in calcium and phosphorous. Mineral is deposited within the interstitium and tubules. Grossly, there are pale streaks and, histologically, basophilic mineral deposits are seen.

Neoplasia
Spontaneous tumors of the urinary tract are uncommon in rats. Tumors reported in the kidney include renal cell adenoma, lipoma/liposarcoma, renal cell carcinoma (with metastasis to the lung and liver), and a nephroblastoma with metastasis to the lung and local lymph nodes. Transitional cell carcinoma of the urinary bladder also occurs spontaneously and in 1 case there was metastasis to the lungs.70,71,76

Obstructive disorders
Uroliths in rats are generally struvite (80.4%) with fewer calcium phosphate (5.9%).19

CHINCHILLAS
Common Problems
Primary renal diseases are poorly described in chinchillas. Urolithiasis has been described, and, although rarely reported, suppurative nephritis is common in younger animals (Fig. 13). E coli has frequently been isolated based in 1 author’s experience (DRR).

Fig. 13. Histology of a suppurative tubular nephritis in a chinchilla (C lanigera). A bacterial culture collected at the time of necropsy isolated E coli. (HE stain, 10 × magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).
Obstructive disorders

Unlike rabbits and guinea pigs, chinchillas eliminate excessive calcium approximately 80% via feces and only 1% to 3% via urine. Nevertheless, their urine calcium concentration may show great individual variation. Urolithiasis is reported in chinchillas and most are calcium carbonate (87.7%) and miscellaneous (6.8%). The clinical signs are as expected, with irritation around the preputial orifice, as well as hematuria, pollakiuria, and stranguria. Most uroliths are found in the urinary bladder, with fewer within the urethra. In males, most are in the proximal urethra near the pelvis and cranial to the os penis (Fig. 14). Prognosis is guarded with urethral stones and recurrence is common. Within the urinary bladder there may be thickening of the bladder wall with an inflamed mucosa.

Perineal hernias are described in 2 sexually intact males. The hernia contents were urinary bladder in 1 animal and lobules of fat in the other. In both cases, the testicles could be moved freely in and out of the inguinal canal.

HEDGEHOGS

Common Problems

Hedgehogs (Atelerix albiventris) have been reported to have cystitis and uroliths, but the most common renal problem is chronic nephritis. As in other species, chronic renal disease is likely to have a large number of causes, such as infections, immune-mediated disorders, and nephrotoxins. Renal disease may also be a part of a systemic condition. Grossly and histologically, the lesion is similar to others that have been previously described in other animals.

Inflammatory/infectious

Cystitis is uncommon in hedgehogs, with scarce descriptions in the literature. In females, uterine tumors are a much more common cause of hematuria. However, 1 author (DRR) has diagnosed cases of hemorrhagic acute to subacute cystitis. Lesions of the urinary system collected from the database of 1 author (DRR) are summarized in Table 1. One case was of encrusted cystitis. There was extensive mineralization of the urinary bladder with an ulcerative and/or necrotic cystitis. This type of lesion is uncommon and has been associated with alkaline urine and several bacteria, including Staphylococcus pseudintermedius and Corynebacterium urealyticum (Fig. 15).

Fig. 14. A chinchilla (C lanigera) with a urinary stone and the os penis can be seen (black arrow). Lateral view. (Courtesy of C. Griffin, DVM, ABVP (avian practice), Kannapolis, NC.)
SUGAR GLIDERS

Common Problems

There are very few reports of urinary disease in sugar gliders.

Inflammatory/infectious
Renal klossiellosis has been described in pet sugar gliders. In affected sugar gliders there is tubular dilatation and interstitial nephritis. The various life cycle stages can be identified within the renal tubular epithelium. Based on morphologic evaluation, the name *Klossiella dulcis n.* sp was proposed. No treatment has been described.\(^\text{82}\)

Obstructive disorders
Bilateral hydronephrosis in a male sugar glider developed secondarily to a functional obstruction suspected to be from hyperplasia and inflammation of the urinary bladder and ureteral epithelium. The animal presented with acute caudal abdominal swelling.\(^\text{83}\)

One author (AML) has noted several cases of disease of the cloacal and paracloacal glands that resulted in apparent urinary obstruction; obstruction resolved with treatment of the primary cause. Both sexes have paracloacal glands (anterior, dorsal,}

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**Table 1**
Data from Zoo/Exotic Pathology Service of 85 hedgehog (*Atelerix albiventris*) urinary lesions from 878 case submissions 1998 to 2019

| Lesion Category | Lesion                           | Number of Hedgehogs |
|-----------------|----------------------------------|---------------------|
| Inflammatory    | Nephritis, chronic fibrosing interstitial | 46                  |
|                 | Cystitis                          | 7                   |
|                 | Cortical infarct                  | 6                   |
|                 | Nephritis, interstitial           | 5                   |
|                 | Glomerulitis                      | 4                   |
|                 | Nephritis, suppurative            | 2                   |
| Neoplastic      | Lymphoma                          | 3                   |
|                 | Sarcoma, renal (not differentiated)| 1                   |
| Miscellaneous   | Renal mineralization              | 9                   |
|                 | Renal tubular necrosis (nephrosis)| 8                   |
|                 | Nephrosis pigmentary              | 3                   |
|                 | Renal cysts                        | 3                   |

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Fig. 15. Hedgehog (*A. albiventris*) encrusted cystitis (HE stain, 20× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).
and ventral), which are more developed in the male. Under the influence of testosterone, the dorsal paracloacal glands and testes increase in size.\textsuperscript{1} Inflammation and neoplastic disease of these glands can result in functional obstruction of the digestive and urinary tract.\textsuperscript{84} There are reports of a paracloacal cyst,\textsuperscript{85} sebaceous nodular hyperplasia, and carcinomas in male sugar gliders. Self-mutilation, stranguria, and percloacal swelling are the typical clinical signs. One case had a dorsal paracloacal gland carcinoma\textsuperscript{84} and the other a transitional cell carcinoma with squamous differentiation.\textsuperscript{86}

**Toxic exposure**

Copper toxicosis has been described in a sugar glider voiding red-brown urine and showing possible seizurelike activity. The kidneys were enlarged and dark red with marked, acute, multifocal renal tubular degeneration and necrosis with hemoglobin casts. Icterus was noted and the liver had marked, acute, centrilobular hepatocellular degeneration and necrosis with hepatocellular copper accumulation confirmed with rubeanic stain and a hepatic copper concentration of 709 parts per million (wet-weight basis). The provided mineral and vitamin supplement had 2.26 and 250 μg/mL respectively. It was unknown whether the vitamin supplement could have led to the toxicosis.\textsuperscript{87}

In a review of sugar glider case submissions (DRR) from 1998 to 2019, out of 337 there were 86 urinary tract lesions. Interstitial nephritis was common, as was the nonspecific finding of renal tubular necrosis (nephritis). Other lesions included membranogluomerulopathy, renal cysts, and transitional cell carcinoma of the urinary bladder (Fig. 16). Lesions are summarized in Table 2.

**TREATMENT OF RENAL INSUFFICIENCY AND FAILURE**

Without benefit of evidence-based therapies for exotic mammals, treatment of renal insufficiency and failure is based on therapies considered most advantageous in traditional pet species.

![Fig. 16. Transitional cell carcinoma in the urinary bladder of a sugar glider (P breviceps) (HE stain, 10× magnification). (Courtesy of D. R. Reavill, DVM, DABVP (Avian and Reptile & Amphibian Practice), DACVP, Carmichael, CA).](image-url)
Chronic Kidney Disease

The 2016 International Society of Feline Medicine guidelines for management of chronic kidney disease (CKD) may be useful. Even in cats, simple and accurate markers for renal function are currently unavailable; biomarkers such as blood urea nitrogen (BUN), creatinine, and urine specific gravity are useful, but interpretation can be difficult. The principles of treatment are supportive, and the overall goal is maintenance of quality of life. Even the goal of maintaining hydration has not been formally associated with increased longevity and improved quality of life in cats with CKD; however, the benefits of normohydration are considered obvious enough to continue recommendation of fluid therapy.88

Unstable or decompensated patients benefit from hospitalization and intravenous (IV) fluid therapy. Fluid rates are based on calculation of hydration deficit: body weight (kg) times estimated dehydration (%) times 1000. Hydration deficit is added to maintenance fluids (50 mL/kg/24 h in cats) and provided over 24 to 48 hours. Once normohydration is achieved, fluid therapy continues at maintenance rate, monitoring the patient carefully for evidence of fluid overload. Fluid therapy is tapered over several days once condition and appropriate biomarkers have improved.

For long-term maintenance of hydration, many recommendations for cats are appropriate in exotic mammals, and include free access to high-quality water using multiple water sources. In exotic pets, this means providing water in bowls along with traditional water bottles. Although moist canned food is not available for rabbits and rodents, moistening greens and hay may aid water intake, as well as supplementing with liquid foods designed for convalescing exotic pets; for example, Oxbow Critical Care or Carnivore Care depending on the nutritional needs of the species. In cats, placement of feeding tubes for oral hydration may be used long term; this approach is not useful for exotic species. However, subcutaneous fluid administration may have utility, because some owners can be taught to administer fluids to small patients at home. Commercial indwelling subcutaneous catheters have been used in cats, and 1 author (AML) has used one in a large rabbit with some success.

Acute Renal Failure

Acute renal failure (ARF) may be reversible if identified and treated early and aggressively. As in other species, ARF may be prerenal (shock, hypovolemia), renal (renal...
disease), or postrenal (obstruction). Urine specific gravity plus changes in BUN and creatinine levels can be helpful. ARF is treated by correcting perfusion deficits (discussed later) and dehydration, and diuresis, along with correction of an identified underlying cause. Once the patient is rehydrated and normotensive, continue fluid administration as maintenance (3–4 mL/kg/h) plus estimated losses (diarrhea/vomiting, if present, and estimated urine volume). Based on information in domestic species, patients with ARF may become 3% to 5% dehydrated each day because of ongoing losses. Fluid administration is decreased when the patient maintains hydration, begins to eat and drink, and BUN/creatinine levels normalize.

IV catheterization is a standard procedure in rabbit medicine; IV access is also possible in large guinea pigs and some chinchillas. IV access may not be possible in any patient that is hypovolemic. For these cases, intraosseous (IO) catheterization is indicated. Placement sites in mammals include the proximal humerus, femur, and tibia.

Placement is best performed in a sedated patient with parenteral analgesia, and a local block (2–3 mg/kg lidocaine into the skin, subcutaneous tissue, and periosteum). Although short spinal needles are useful in rabbits and large rodents, most IO catheters are simple hypodermic needles, 25 to 18 gauge depending on the size of the bone cavity. Seat the needle into the proximal bone, and then orient the needle into the bone cavity. Cover with an IV catheter cap and secure to the limb with tape (Fig. 17).

Treatment of shock in rabbits and rodents has been described. Management of fluid deficits is maintenance fluids estimated at 3 to 4 mL/kg/h in exotic mammals, or via allometric calculations of resting energy requirements (weight [kg]$^{75} \times 70$) plus hydration deficit, adjusting fluid rates for ongoing losses, such as polyuria in cases of renal insufficiency.

Other therapies include erythropoietin for treatment of anemia, analgesia as indicated, and support feeding.

Fig. 17. An intraosseous catheter in place for fluid diuresis for a pet rat (R norvegicus). A 25-gauge needle was used for the catheter, which was taped into place and capped with an IV catheter cap. (Courtesy of A. Lennox, DVM, DABVP (Avian Practice, Exotic Companion Mammal Practice), Diplomate ECZM (Small Mammal), Indianapolis, IN).

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