Case Report — Gevalverslag

Vulvovaginectomy and neo-urethrostomy for treatment of haemangiosarcoma of the vulva and vagina

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ABSTRACT

Vulvovaginectomy and neo-urethrostomy were performed in a 9-year-old German shepherd dog following a diagnosis of infiltrative vulvar and vestibulovaginal haemangiosarcoma. The dog was presented for intermittent vulvar haemorrhage over a 3-month period. On examination the vulva and vestibulovagina were distended and firm. Vaginal discharge and fine needle aspiration cytology detected anaplastic cells. Haemangiosarcoma was diagnosed on biopsy. A new urethral opening was created in the floor of the vagina allowing resection of the vulva and caudal vestibulovagina. Urinary continence was preserved and healing was without complications.

Key words: urethrostomy, vaginal, vulvar, vulvovaginectomy.


INTRODUCTION

Primary tumours of the vulva and vagina account for 2–3% of neoplasms in dogs. Most are benign, with leiomyomas and fibromas being most common. Leiomyosarcoma and carcinoma are frequently-found malignant tumours, whereas the transmissible venereal tumour (TVT) is a more common tumour, especially in resource-poor communities in South Africa.

Tumor resection, commonly facilitated by dorsal episiotomy, is successfully used to treat benign tumours. Vulvectomy for malignant tumours has been described, but tumour recurrence is common, and patients with tumours deemed inoperable are often euthanased. More extensive vulvovaginectomy and perineal urethrostomy has been described for infiltrative tumours of this region. Haemangiosarcoma of the vulva and vestibulovagina in the dog is rare, with only a single case reported previously.

The purpose of this report is to describe the clinical manifestations of a genital tract haemangiosarcoma and surgical treatment performed, as compared to previously published descriptions of treatment of malignant tumours in the vulval area.

Fig. 1: Enlarged distended vulva and caudal vagina. There is also a surgical wound on the left-hand side, which represents the initial biopsy site.
normal on palpation. Urination produced a strong urine flow with blood clots present in the initial part of the urine stream.

On digital examination, the vestibular and caudal vaginal lumen was very narrow as a result of a large intramural mass compressing the vaginal wall. The mass was approximately 7 cm in size and occupied the entire floor and right wall of the vulva, vestibulum and caudal vagina. The narrow vaginal lumen and ongoing haemorrhage hindered vaginoscopic examination. Numerous blood clots and a large erythematous area were observed on the vestibular floor. Multiple areas of petechiae and ecchymoses were present throughout the vestibular and caudal vaginal mucosa as well as small ulcerations over the mass. The urethral opening could not be visualised.

Haematology revealed marked leucocytosis (30.1 × 10^9, normal 6-15), mature neutrophilia (24.98 × 10^9, normal 3.0-11.5) and monocytosis (1.51 × 10^9, normal 0.15-1.35). Serum chemistry was unremarkable. A urine sample collected by cystocentesis appeared normal macroscopically and had a pH of 7 and specific gravity of 1.025. Moderate proteinuria, haemoglobin crystalluria and few erythrocytes were detected on urinalysis. Smears obtained from the vulvar discharge for cytological examination were highly cellular, with obvious secondary infection and numerous anaplastic cells. However, no specific tumour diagnosis could be made. Fine needle aspirates of the mass demonstrated solitary and aggregated anaplastic cells. A presumptive diagnosis of anaplastic sarcoma was made.

Thoracic radiographs for lung metastases revealed no evidence of atrial metastases. The perineal area was surgically prepared in a standard manner and included a purse-string suture of the anus. The animal was placed in sternal recumbency with the perineum elevated. The urethral opening was identified following a dorsal episiotomy. It was displaced dorsolaterally to the right and recessed within the tumour (Figs 2, 3). A 16F Foley catheter (Macmed) was placed to drain the bladder and identify the urethra. A fusiform

![](image1)

Fig. 2: Dorsal episiotomy – the external urethral is catheterised (curved arrow) and displaced by the haemangiosarcoma (arrowhead).

![](image2)

Fig. 3: Schematic presentation of surgical excision. a: Episiotomy to visualise tumour and urethral opening; b: fusiform incision to include vulva, episiotomy and tumour; c: exteriorised vulva and vagina, urethroscopy over catheter; d: position of urethroscopy in floor of vagina.

An incisional biopsy was taken from the left lateral aspect of the vaginal mass. Areas of necrosis, fibrosis, haemorrhage and neutrophil infiltration were found on histopathological examination. Nests of anaplastic cells with a high mitotic index were found in the necrotic areas and in some instances formed ill-defined vascular channels. A diagnosis of anaplastic, infiltrative haemangiosarcoma of the vagina was made. Echocardiography revealed no evidence of atrial metastases.

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skin incision was made around the vulva, vestibulum and vagina so as to encompass the ventral vulval commissure, the palpable portion of the vestibulum and vagina dorsally, the previous biopsy site and the episiotomy incision. The constrictor muscle of the vestibulum was transected below the junction with the external anal sphincter. The ischiocavernosus and ischiourethralis muscles were detatched from the ischial rim. Branches of the ventral perineal artery were ligated or electrocoagulated as encountered.

The cranial extent of the tumour was identified through the episiotomy incision. The Foley catheter/urethra was palpated and identified. A 2 cm longitudinal incision was made over the catheter 2 cm cranial to edge of the tumour. The longitudinal incision extended through the vaginal mucosa, vaginal and urethral muscle and urethral mucosa to expose the intramural intrapelvic portion of the urethra. The urethral mucosa was sutured to the vaginal mucosa with 5/0 simple interrupted polydioxanone (PDSII, Schering-Plough) sutures and a new urethral opening created. The vulva, vestibulum, caudal vagina and distal part of the urethra, together with the tumour were resected immediately caudal to the neourethrostomy site. A small amount of tumour tissue was identified within the vaginal muscle layer on the left periphery. An additional 2 cm section of the vagina was resected obliquely on the left. Subcutaneous sutures using 30 polydioxanone were placed to eliminate dead space and eliminate skin tension. The skin was sutured to the muscular and mucosal layers of the vagina using 4/0 polydioxanone in a simple interrupted pattern (Fig. 4).

Histopathological examination of the excised tumour confirmed the incisional biopsy results. Further immunoperoxidase staining was positive for Von Willebrand factor, confirming a haemangiosarcoma. Although the lateral margins of the surgical field were free from neoplastic cells, the deeper portion of the surgical excision contained neoplastic cells.

Post-operatively, the dog was given 0.01 mg/kg buprenorphine (Temgesic, Schering-Plough), repeated 8 hours later. Thereafter, ketoprofen (Orucote, Rhône-Poulenc Rorer) at 1 mg/kg once a day for 2 days and amoxycillin/clavulanic acid (Synulox, Pfizer) at 20 mg/kg twice a day for 10 days were administered. Petroleum jelly was appliedtopically around the new vaginal opening. A strong urine flow was present but urination was uncomfortable for the dog for the first 24 hours. No signs of urinary incontinence were noted. Further post-surgical recovery was uneventful. The owner declined adjunct chemotherapy with doxorubicin.

Five months post-operatively, signs of vaginal bleeding returned and tumour recurrence was diagnosed on clinical examination. The dog was euthanased at the request of the owner. No post mortem examination was performed.

DISCUSSION

Primary tumours of the vulva and vagina are usually (71–82%) benign2,7. In a report by Kydd and Burnie5, who examined 21 cases of vulvar and/or vaginal neoplasia histopathologically, 20 of the 21 cases were found to be benign. Apart from leiomyomas and fibromas, other types of benign neoplasia reported are lipoma, histiocytoma, benign melanoma and myxoma. Malignancy accounts for only a small percentage of vulvar and/or vaginal tumours, particularly if TVT is performed. Only single cases of breast tumour, epidermoid carcinoma, haemangiosarcoma, osteosarcoma and adenocarcinoma have been reported2,7. Magné and others7 reviewed 7 cases that presented with a vaginal and/or vestibular mass that resulted from a caudal extension of a bladder or urethral carcinoma.

The most common clinical signs of vulval/vaginal tumours are an abnormal vulval discharge or the sudden appearance of a protruding mass. Many of the benign tumours are pedunculated, grow within the confines of the vagina and only at a later stage protrude through the vulval lips7. Leiomyomas originate from the smooth muscle of the vagina or vestibulum and may be intramural or extraluminal6. Other presenting signs may be vulval or perineal swelling, dysuria and tenesmus. Urinary incontinence does not occur8. The average age on presentation depends on the type of tumour. In general, dogs with TVT present at younger age (average age of 4.4 years), whereas dogs with other tumours present at a much older age (10.8–11.2 years)2,5,8.

Most malignant tumours are non-pedunculated1. Vulvectomy has been described as a treatment for malignant tumours, but recurrence is common7. In some cases, tumours have been deemed inoperable following exploratory episiotomy, and the animal subsequently euthanased1. The average survival time is approximately 30% shorter than those with benign tumours7. A more extensive surgical resection technique combining
vulvovaginectomy and perineal urethrostomy has been described for large and infiltrative tumours of this region. This case of vaginal haemangiosarcoma in the dog is only the second published case. In the previously-reported case, the tumour was incompletely excised with rapid regrowth of the tumour within 1 month. Although the tumour described in this report was incompletely excised, tumour regrowth occurred only 3 months post surgery.

The surgical technique described here avoids creating a perineal urethrostomy, but rather creates a neo-urethrostomy site within the remaining proximal portion of the vagina. Deep dissection of the pelvic urethra with possible damage to the nearby caudal vesicular artery, ureters, pelvic plexus and pudendal nerve branches is thus avoided. It is possible that the undisturbed longitudinal muscle fibres of the vaginal and urethral muscle immediately surrounding the new urethrostomy site may be helpful in compressing the opening and preventing retrograde urethritis and cystitis. Sutures between the vaginal and urethral mucosa were without tension. Suturing the urethra directly to skin requires caudal traction of the urethra and is more likely to result in tension on the suture line. Excessive suture line tension resulting in wound dehiscence is a common cause of urethral stricture formation in perineal urethrostomies in cats. A disparity in the thickness of the tissue edges was not experienced.

With a neo-urethrostomy, urine is directed onto vaginal mucosa rather than skin, which may reduce problems with urine scald. The remaining portion of the vagina, cervix and uterus remains connected to the outside. Also, the peritoneal cavity is not entered. Dissection of the urethral stump or removal of the uterine remnants may cause penetration of the peritoneal cavity and allow seeding of neoplastic cells or infection into the abdomen. It is also possible that urovagina may occur if the neo-urethrostomy opening lies dorsal to the ischial arch. To prevent this possible complication the neo-urethrostomy opening must be caudal to the ischial arch.

Vulvovaginectomy and neo-urethrostomy may be curative for large benign tumours involving the vulva and vagina of the bitch with minimal post surgical complications such as urinary incontinence, retrograde urethritis and urine scald. A similar result may be possible with early malignant tumours but palliation may still be achieved for advanced malignancies, as was shown in this case.

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REFERENCES