Vaginal paraurethral leiomyoma
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ABSTRACT
Leiomyomas of the vagina are uncommon soft tissue tumors. Only a few cases are reported in medical literature. It is usually seen between 20-80 years of age. We report a case of 30-year-old woman with paraurethral leiomyoma. Leiomyomas in this location should be differentiated from urethral diverticula and vaginal cysts. Complete surgical excision should be the choice of treatment. Excision usually has good prognosis and rarely recurs.

Keywords: Leiomyoma, Vagina, Paraurethral, Surgery

INTRODUCTION
Leiomyomas develop most commonly in the uterine corpus and much less often in the cervix. They may develop in round ligament but this is rare. Growth of leiomyomas is dependent on estrogen production. Tumor thrives during years of highest ovarian activity. Leiomyomas are common tumours of mesenchymal origin, well-circumscribed and can develop where smooth muscle is present.1 Leiomyoma outside uterus has been reported in various sites such as vulva, perineum and buccal mucosa.2 There are few reports of leiomyomas arising from paraurethral region.3 There can be diagnostic difficulty and hence we are presenting this case.

CASE REPORT
A 30 year old married woman, gravid two presented to out-patient clinic with complaints of small swelling over vulva since 3 years. Since 15 days she noticed sudden increase in size causing discomfort. There was no history of urinary complain, sexually transmitted disease, fever or trauma. General examination revealed a healthy woman. On local examination there was 6x7 cm mass arising from anterior vaginal wall near urethral orifice on right side at 10 o’clock position (Figure 1 & 2).

Mass had smooth surface with few areas of slough. On palpation it was mobile and minimally tender. Ultrasound examination of pelvis was normal. Under spinal anesthesia, urethroscopy and biopsy was taken which was normal. Excision of the mass was planned. Foley’s catheter was inserted and urethral opening was shifted to one side. Incision was taken medial to labia minora. The tumors got separated from periurethral tissue and was dissected from the surrounding tissue. The dissection in the paraurethral region was meticulous to reduce injury to the urethra. Complete excisional biopsy of the mass was performed. Minimal bleeding was encountered during excision. A two-layer closure was performed.

There was no postoperative complication and she gained normal lower urinary tract function after the removal of catheter 24 hours later. At 2 months follow-up, clinical examination showed normal vaginal anatomy. Patient is on follow up and is free of recurrence after a year.
The excised mass was 6x7 cm diameter, firm in consistency, cut surface was smooth whitish, no hemorrhage and necrosis. Histopathology showed well-differentiated spindle cells arranged in orderly intersecting fascicles. These cells had eosinophilic cytoplasm and mostly bland, uniform, cigar-shaped nuclei, resembling normal smooth muscle cells (Figure 3). Immunohistochemistry was positive for muscle markers including smooth muscle actin and desmin (Figure 4).
DISCUSSION

Leiomyoma is a benign neoplasm commonly seen in myometrium of uterus. It is hormone sensitive tumour. It develops during hormonally active reproductive years and regresses after menopause. It originates from smooth muscle cells. It has been hypothesized that origin of leiomyoma is from smooth muscle cells of myometrium due to somatic mutation. There is progressive loss of growth regulation. In previous literature numerous extraterine sites of leiomyoma have been described. There are few reports of paraurethral leiomyoma. Clinically it is difficult to know nature of these tumours.

In this case patient noticed small asymptomatic lump which grew rapidly in two weeks. Patient can present with profuse bleeding. Urinary symptoms are rarely described. Occasionally Paraurethral leiomyoma may cause urinary obstruction. The distinction among urethral, paraurethral, and anterior vaginal wall leiomyoma can be very difficult owing to their anatomic proximity.

The differential diagnosis includes cystocele, urethrocele, Gartner’s duct cyst, urethral diverticula, vaginal cyst, Bartholin gland cyst and vaginal malignancy.

Transabdominal and or transvaginal ultrasonography and MRI are helpful in establishing the morphology and relationship with adjacent anatomic structures although these tests are not mandatory pre-operatively.

Surgical enucleation is the treatment of choice via vaginal approach. In our case the mass was superficial and easily accessible hence surgery was done with minimal morbidity. However large giant masses greater than 10cms have been removed transvaginally and transperitoneally. Preoperative embolization of a large vaginal leiomyoma to reduce intra-operative blood loss has been described. The procedure is indicated in large hyper-vascularized tumours presenting with haemorrhage. Though benign and slow growing vaginal leiomyoma may occasionally recur. Hence patient should be followed up for recurrence.

Slides advocate the use of immunohistochemistry (IHC) to establish the exact diagnosis. Immunohistochemistry (IHC) for Smooth Muscle Actin (SMA) showed diffuse intense positivity in vascular wall and intervascular stroma confirming their smooth muscle nature (Figure 3 and 4). Patient is on follow up and is free of recurrence after a year.

CONCLUSION

Leiomyomas in extraterine sites present a diagnostic challenge due to their rarity. Perineum, vulva, ovaries, urinary bladder, and urethra can be some of sites for these tumours. Ultrasonography might be of help in establishing the diagnosis. Leiomyomas are treated with conservative surgery. After the surgery long-term follow up is advised. To conclude, this rare case of vulval leiomyoma is reported for its rarity in literature and to bring the awareness of vaginal periurethral leiomyomas.

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