A large cellular angiofibroma of the male pelvis presenting with obstructive voiding: A case report and review of the literature

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Abstract

Cellular angiofibromas (CAF) are rare, benign soft-tissue tumours. The diagnosis of CAF is important given the heavy resemblance to other tumours. Herein, we describe a case of a rapidly growing, very large (13.5 cm) CAF located in the deep pelvis of a middle-aged male who presented with difficulty voiding.

Introduction

Cellular angiofibromas (CAF) are rare, benign soft-tissue tumours first described by Nucci and colleagues in 1997.1 CAF share pathological features of angiomyofibrolastoma and aggressive angiomyxoma (AAM), but do not fall into either category. The masses are well-circumscribed, typically less than 3.0 cm and show marked female predominance. In 1998, Laskin and colleagues reported 11 of these tumours in male genitalia,2 and since then there have been only a few case reports. Most have been found in women, primarily in the vulva.3,4 In males, most are found in the scrotum and inguinal region, averaging about 5 to 6 cm in size.2,3 The diagnosis of CAF is important given the heavy resemblance to other tumours, such as AAM, which is known to have a very high rate of local recurrence.4 Herein, we describe the case of a rapidly growing, very large (13.5 cm) CAF located in the deep pelvis of a middle-aged male who presented with difficulty voiding.

Case presentation

A 58-year-old man initially presented with right groin/pelvic pain radiating down into his scrotum, as well as decreased urinary flow and decreased ejaculation. Due to refractory symptoms, a computed tomography (CT) scan of the abdomen and pelvis was done which showed a 7 × 7 × 5-cm mass to the right of the prostate extending down to the bulbar urethra. Prior attempts at both transurethral drainage and transperineal aspiration were unsuccessful and he was then referred to our clinic. A pelvic magnetic resonance imaging (MRI) revealed substantial growth; a 11-cm well-circumscribed lesion abutting the rectum and displacing the urinary bladder and prostate proximally (Fig. 1, Fig. 2). Biopsies showed a benign stromal tumour of genitourinary origin. Resection was recommended via combined abdomi-perineal approach with colorectal surgery and urology.

Exploratory laparotomy showed a large encapsulated mass occupying the entirety of the lower pelvis. The mass was dissected off of the prostate with relative ease, but was more difficult distally owing to dense adherence to a 3-cm portion on the anterior surface of the bulbar urethra. The mass was carefully freed from the rectum through the peri- neal incision. Ultimately, the intact mass in its entirety was removed with no evidence of residual disease. The specimen was sent to pathology and was read as a large benign spindle cell tumour mostly consistent with a CAF.

Gross description

This was an encapsulated, well-circumscribed mass with a thin fibrous capsule. It was elongated and measured 13.5 × 7.4 × 3.7 cm (Fig. 3). The mass felt rubbery and sectioning revealed a gelatinous, tan-pink to grey surface that was focally trabeculated.

Microscopic description

The mass consisted of loosely arranged bland spindle shaped cells in a myxoid stroma (Fig. 4). Abundant partially hyalinized vessels were interspersed throughout the lesion. Dense collagen bundles, as well as thin filaments, were also evident. There was no significant pleomorphism.
Discussion

CAFs are rare benign mesenchymal tumours normally found in women; it is rarely found in men. The median age at diagnosis is the mid 50s, with a size typically about 5 to 6 cm. There are only 2 reported cases measuring more than 10 cm. CAFs are mainly found in the scrotum and inguinal region, but have also been reported in the knee, upper eyelid, oral and perianal region. Grossly, they are well-circumscribed superficial lesions with a thin fibrous capsule. Histopathologically, they are moderately cellular with prominent hyalinized round vessels and a uniform distribution of bland spindle cells. Approximately 60% express CD 34; 35% to 55% express estrogen receptor/progesterone receptor (ER/PR) and 10% express desmin. S-100 is uniformly non-reactive.

Our mass showed immunoreactivity for CD34, desmin and ER/PR, but was non-reactive for S100, smooth muscle actin, pan keratin and CD117. This case is unique as it is one of the first cases of CAF found deep in the pelvis, extending to the perineum, abutting the prostate and rectum, and densely adherent to the urethra. The fact that it was about 13.5 cm on excision also shows that tumours in this location can grow as large as any previously reported CAF. This tumour also underwent rapid growth, expanding by almost 50% over a few months. This has not been previously described and bolsters the current practice of surgical excision.
We present a large CAF of the male deep pelvis found with the workup of refractory obstructive voiding along with pelvic pain. The diagnosis is made with imaging showing a well-circumscribed hypervascular mass containing fat; histology showing prominent hyalinized round vessels and a uniform distribution of bland spindle cells, and gross exam showing an encapsulated rubbery mass. Given the rapid growth and lack of recurrence or metastasis, simple surgical excision is the treatment of choice for symptomatic lesions.

**Competing interests:** None declared.

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**References**


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