Recurrence of aggressive angiomyxoma of the vulva: a case report

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ABSTRACT
Aggressive angiomyxoma (AA) of the vulva is a rare mesenchymal neoplasm that typically affects women of childbearing age, with a high rate of local recurrence. We present the case of a 38-year-old woman with local recurrence of aggressive angiomyxoma of the vulva 5 years after surgical excision of the first lesion. Aggressive angiomyxoma is diagnosed by histopathological and immunohistochemical features. The therapy is surgical, followed by a long follow-up or hormonal therapy to prevent recurrence.

Key words: Angiomyxoma; vulvar neoplasms; mesenchimal tumor; case report; local recurrence

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INTRODUCTION

Aggressive angiomyxoma (AA) is a rare onset mesenchymal neoplasm that affects the pelvic and perineal region in reproductive age. It most frequently affects women in the fourth decade of life. It is a benign neoplasm first described in 1983 by Steeper and Rosai (1). It is characterized by a slow increase in volume, a high rate of local recurrence and a low tendency to metastasize (2).

CASE PRESENTATION

A 38-year-old woman, G1 P1 (one cesarean section), comes to our observation following the onset of vulvar neoformation on the large left lip of about 5 cm in diameter. The lesion appears of a soft consistency, movable on the superficial and deep planes and covered by apparently healthy skin. The patient reports, in her anamnesis, about 5 years ago, excision of cysts of the left Bartolini's gland with histological outcome of "aggressive deep angiomyxoma". Performs MRI of the pelvic excavation which highlights "at the level of the large left lip, presence of an elongated aspect extending in an antero-posterior direction for about 6 cm, with maximum thickness at anterior level of 14 mm with significant contrast enhancement after administration of contrast agent. The patient, with suspected recurrence of aggressive angiomyxoma of the vulva, is therefore subjected to surgical exeresis of the vulvar neoformation (figures 1-3). The histological examination confirms the diagnosis of aggressive deep angiomyxoma, with immuno-histochemistry of desmin positive neoplastic cells, smooth muscle actin (SMA), estrogen receptors, progesterone, CD34. The margins of resection are undamaged. It is therefore included in a six-year follow-up program for 5 years.

Figure 1-2. Surgical excision of the vulvar AA.
DISCUSSION

Aggressive angiomyxoma is a benign mesenchymal tumor that affects almost exclusively the genital, perineal and especially the vulva of women of reproductive age. It is characterized by slow growth but a high rate of recurrence even after several years after the first diagnosis (3). AA is mistakenly diagnosed in 80% of cases. Clinically it is, in fact, initially recognized as a cyst of the Bartolini’s gland, lipoma, cyst of the small or big vulvar lip, Gartner duct cyst, etc. It enters in differential diagnosis also with neofor- mations such as superficial angiomyxoma, angiomyofibroblastoma, cellular angiofibroma, leio-myofibroma and any polypoid neoformation of the perineum. In fact, it appears as a soft neoformation, such as to make the real evaluation of its volume difficult because of its consistency and its tendency to deeply invade the pelvic tissues, covered by healthy skin. It can be symptomatic and present with vulvodynia, dysuria, sense of weight or occasionally found during a gynecological check-up (4). Therefore the clinical diagnosis is very complex due to the rarity of this formation and the symptomatology common to other lesions of the genital sphere which sometimes appears shaded or completely absent. For this reason it is possible to use different diagnostic methods for images that include ultrasound, Computed Tomography and Magnetic Resonance. Among these, Magnetic Resonance (MRI) plays a greater role due to the greater amount of information it provides (5). Indeed, MRI shows the isointense lesion in T1-weighted and hyperintense images in T2-weighted images due to the high-water content and the presence of free AA-typical mixoid matrix. It also allows a precise evaluation of the overall volume of the new formation and its limits with respect to the surrounding structures (6). The diagnosis is given by a set of histopathological features such as: stellate and spindle cells with poorly defined cytoplasmic margins and separated by abundant myxoid stroma and fibrillar collagen. The AA is also characterized by a well-represented vascular component with many thin vessels. There is no evidence of atypical mitotic activity or cellular atypia (7). Immunohystrochemistry of the lesion is characterized by receptors positive for estrogens, progesterone, vimentin, desmin, SMA, CD34 and CD44 and always negative for S-100, carcinoembryonic antigen (CEA) and keratin. There is no consensus regarding the pathogenesis of this lesion. Recent cytogenetic studies have shown the presence of genetic alterations affecting chromosome 12, in the 12q13-15 region. In this region there is the HMHI-C gene (high mobility group protein isoform C) that encodes proteins involved in the regulation of gene transcription involved in the pathogenesis of AA. The possibility of using mutated anti-HMHI-C antibodies could be used to identify microscopic residue of neoplasia (8).

CONCLUSIONS

Therapy in the first instance is therefore excisional surgery, followed by follow-up or hormonal therapy. The hormonal treatment based on tamoxifen, raloxifene or Gn-RH analogues finds space in case of partially excised bulky lesions or in the treatment of relapses. Chemothary and radiotherapy are not useful in the treatment of these lesions due to the poor mitotic activity of the AA (9). Finally, it is im
important to keep in mind that AA is characterized by a high rate of recurrence that can occur even after many years, so it is important to set a follow-up period of not less than 5 years.

CONFLICT OF INTERESTS

The authors declare that they have no conflict of interests.
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