Recurrent Aggressive Angiomyxoma

Angiomioksma Agresif Rekurens

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ABSTRACT

Aggressive angiomyxoma (AA), a rare mesenchymal tumor with high recurrence rate and infiltrative nature, most generally arises in the vulvovaginal region, perineum, and pelvis of women in reproductive age peaking from age 31 to 35 but uncommonly found on men. It appears as a slow-growing gelatinous mass, which commonly asymptomatic. Imaging studies including USG, CT scan, and MR imaging are of great importance for the diagnosis of AA on identifying the mass characteristics. This paper reports a case of a 37 year old P3A0 woman with a lump on the left side of vaginal lip for 2 weeks before admission. The lump was soft to touch and was not painful, without other accompanying symptoms. Past medical history showed the lump has recurred after being surgically removed. On physical examination, a lump was found on the left vulva with soft consistency. CT scan showed a well-circumscribed hypodense mass on left labia majora and showed post contrast wall enhancement. Fine needle aspiration biopsy was done and showed hypocellular smear with myxoid matrix and bleeding. The mass was then removed by vulvectomy procedure. Histopathological analysis of the mass shows invasion of myxoid stroma with blood vessels consistent with aggressive angiomyxoma. In conclusion, AA is usually asymptomatic and has high tendency of recurrence. Imaging procedure is useful on identifying mass characteristics and biopsy can be done to establish AA diagnosis. Surgery is the main modality for the treatment of AA, and administration of gonadotropin-releasing hormone analogue can prevent further recurrence.

Keywords: Aggressive angiomyxoma, gonadotropin-releasing hormone agonist, recurrent

ABSTRAK


Kata Kunci: Angiomiksoma agresif, gonadotropin-releasing hormone agonist, rekurens
INTRODUCTION

Aggressive angiomyxoma is a rare tumor of vulvovaginal and perineal region. Although the tendency to infiltrate adjacent organs emphasize its aggressiveness, it is classified as benign tumor since it has no metastasize capability (1). The incidence of this disease peaks on the third decade of life and most commonly occurred on female (2,3). This disease has very high recurrence rate after being surgically removed (4) and is accompanied with scarce symptoms occurrence (4), thus every patient with recurrent painless nodule in perineal and vulvovaginal region has to pay more attention towards this disease.

In this paper, we report a case of aggressive angiomyxoma with multiple prior recurrences with history of complicated surgery by a general surgeon, and resolved by gynecologist oncologist's surgical intervention. This case represents a most typical appearance of aggressive angiomyxoma considering its recurrence and asymptomatic nature, and the significance of gynecologist intervention, especially oncologist gynecologist, on management of this patient.

CASE REPORT

We reported a case of a 37 year old female with lump on the left side of her vaginal lip for 2 weeks before admission. She admitted the lump on the left side of her vaginal lip was suddenly getting bigger. The lump was soft to touch and was not painful. She denied whether having fever, abnormal vaginal discharge, or bleeding from her genitalia. She also denied having difficulties in passing motion and urinating. Pelvic examination revealed a 5x3 cm lump with soft consistency on left vulva. Cervical and vaginal examination turned out to be normal.

Previously, the patient admitted she had a lump on the left side of her vaginal lip 3 years before admission. In the beginning, she found a lump that looked like a boil. She applied some Chinese medicine on top of the lump every day, and the lump burst and released pus on the third day, then the pus resolved. One year before admission, she found the same lump on the left side of her vaginal lip. She applied the same Chinese medicine on top of the lump and it burst and released pus from the lump then the lump resolved again. Few months later, she found same the lump again on the same spot. She applied the same Chinese medicine, but this time it did not work. She consequently chose alternative medicine and took herbal medication for 2 weeks, but she stopped taking those medications because the lump was getting bigger overnight. The lump was very painful and appeared very swollen. She described the appearance that the size was as big as a duck egg. The patient was then hospitalized in Pontianak for 2 weeks and underwent FNAB and CT-scan but with no reassuring result.

Her CT-scan showed circumscribed hypodense mass on left labia majora with 8.5x2.7cm of size with post contrast wall enhancement (as seen in Figure1), while FNAB examination showed hypocellular smear with myxoid matrix and bleeding. Then patient was transferred to Siloam MRCCC Jakarta and hospitalized for 1 month to undergo a surgical procedure by a general surgeon. The surgeon failed to totally excise the mass because of massive bleeding due to cut on a number of blood vessels on the surgical site.

The patient required eight bags of blood transfusion, and admitted to intensive care unit. A severe infection was developed on her genital 3 days after admission. The patient said that the operated area was foul-smelling, bleeding, and pus was found around the suture. The patient also developed high fever thus she received intravenous antibiotics until her general condition improved. The patient was consulted to oncologist gynecologist in Siloam MRCCC and was planned for a vulvectomy and tumor removal with wide excision. The
surgery went well and successfully removed the mass with 2 cm tumor margin, as shown in Figure 2 below.

Figure 2. Aggressive angiomyxoma mass after being surgically removed

Note: Histopathological analysis specimen from labia majora sinistra: Proliferation of blood vessels are found, among collagen tissue and myxoid tissue. Among the myxoid tissue, stellate or spindle cells can be found. No nuclei mitotic or atypia are found. Conclusion: Aggressive Angiomyxoma, malignancy is not found.

DISCUSSION

Aggressive angiomyxoma is a rare mesenchymal tumor that most commonly arises in the vulvovaginal region, perineum, and pelvis of women (1). Aggressive angiomyxoma (AA) is most commonly found in women, but less commonly found in men (2). The age distribution of aggressive angiomyxoma is wide, commonly in reproductive age with the peak incidence from 31 to 35 years old (3). Our rare case also happen in her reproductive age, which was first found at 34 years of age.

The term aggressive was introduced to emphasize the locally aggressive behavior and the high potential for local recurrence (36-72%) (4). It is defined as benign but has infiltrative potential into skeletal muscle and fat. The disease is therefore considered locally aggressive although it does not infiltrate surrounding vissus. The etiology is unknown. Aggressive angiomyxoma (AA) is most often found in or in proximity to the lower pelvis, more specifically perineum, vulva, vagina, or inguinal regions. The majority of patients present with a slow-growing mass, often more than 10 cm in the largest diameter, which is asymptomatic and this is frequently the only symptom that occur, just like what we found in this case. Observed accompanying signs and symptoms are pain around the affected area, feeling of pressure, or dyspareunia. Our patient denied any pain at first. It presented as a painless, poorly circumscribed gelatinous vulvar mass and clinically simulates a Bartholin gland cyst or an inguinal hernia.

The tumor cells are characteristically positive for estrogen and progesterone receptors, suggesting a hormonal role in the development of the tumor (1). Chromosomal translocation of the 12q13-15 band involving the HMGA2 gene has been described. Aggressive angiomyxoma (AA) must be diagnosed considering the age, clinical evolution, location, imaging assessment and anatomopathological and immunohistochemical evidence (5). However, due to the rarity of cases and lack of typical characteristics, the preoperative diagnosis is very difficult, hindering the therapeutic planning. Thus, most cases are confirmed histopathologically after the primary surgical resection (6).

Imaging studies are of great importance for the diagnosis of AA. The USG can reveal a polypoid, hypoechoic mass of soft tissue, which may appear as a cyst. At the CT, the characteristics vary and may include a homogeneous well-defined and hypodense mass relative in relation to the muscle, a hypocattenuating solid mass with an internal spiral pattern after intravenous contrast agent or a predominantly cystic mass with solid components (7). On T2-weighted MR imaging, the tumor has high signal intensity (Figure 3).

Unfortunately, our patient did not undergo the MRI procedure with contrast, which may lead to misdiagnosis this rare angiomyxoma.

On gross examination, the tumors are characteristically soft, bulky masses with a smooth external surface. The histopathological examination shows myxoid stroma, hypocellularity, small spindled and stellate mesenchymal cells with undefined cytoplasm. So, some may underestimate it as a common soft tissue tumor or did not notice it. Pleomorphism and mitoses are not present and there is no evidence of coagulation necrosis in the tumor.
cells. Characteristically, there is a prominent vascular component with vessels of various calibers. Infiltration into fat, muscle, and nerves is seen (9,10). Immunohistochemically, most AAs are positive for desmin, smooth muscle actin, muscle-specific actin, vimentin, oestrogen receptor, and progesterone receptor (4). The differential diagnosis of AA is based on other forms of soft tissue tumors (myxoma, myxoid lipoma, neurofibroma) and malignant tumors with metastatic potential (myxoid liposarcoma, myxofibrosarcoma, embryonal rhabdomyosarcoma) (7). Additionally, it can occasionally mimic Bartholin’s cyst, labial cyst, Gartner’s duct cyst and perineal herniation. The distinction of these tumors can be seen on Table 1.

Because of its high rate of local recurrence, the first line treatment for AA is complete surgical resection with tumor free margin (1). Incomplete lesion resection occurs in 45-66% of cases and is directly associated with the possibility of recurrence, often requiring re-excision (3). Angiographic embolization can be used to adjunct the surgical removal of the tumor mass since it can shrink the tumor and make it easier to identify the tumor from surrounding normal tissues (11). Due to the low mitotic potential of this neoplasm, chemotherapy and radiotherapy are ineffective treatment (12). Treatment with gonadotropin-releasing hormone agonists is an emerging therapy and also can be used in case of incomplete resection or recurrence (1). As what happened in our patient, due to the vascularization and the bigger size of the tumor itself, the excision procedure is sometimes troublesome, but it can be prevented by a holistic approach.

The patient in this report exhibited a characteristics of AA based on the literature. The mass this 37 year old patient has on her vulva was soft and painless with no accompanying symptoms. These findings should raise initial suspicion to AA, however, other differential diagnosis such as lipoma and neurofibroma also have to be considered due to its painless nature. After its recurrence, the mass started to be painful. This is mostly because of intervention of traditional medicine this patient received for her vulvar mass. The CT scan result shows typical AA mass embryonal rhabdomyosarcoma) (7). Additionally, it can occasionally mimic Bartholin’s cyst, labial cyst, Gartner’s duct cyst and perineal herniation. The distinction of these tumors can be seen on Table 1.

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