Case Report

Primary ovarian leiomyoma: a diagnostic dilemma

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ABSTRACT

Among all benign primary ovarian smooth muscle tumors, ovarian leiomyoma is uncommon accounting for .5-1% of all tumors. Most of these are unilateral small in size and generally occur in premenopausal women. This paper presents a case report of primary ovarian leiomyoma i.e. 29 years old women with complaints of abdominal pain. The right ovarian mass was histologically diagnosed as ovarian leiomyoma. Unlike other smooth muscle tumors of uterine origin, there are no definite histological criteria to categorize smooth muscle tumor of ovary.

Keywords: Leiomyoma, Magnetic resonance imaging, Ovarian tumor

INTRODUCTION

Primary ovarian leiomyoma is a rare benign tumor of ovary seen in women between 20-65 years. It accounts for 0.5-1% of all benign ovarian tumor.1

Ovarian leiomyoma is particularly unilateral and small but they may reach big size upto 11 kg. It usually occurs in premenopausal women most of these tumors are discovered incidentally.2

Patients are usually asymptomatic, and the tumor is most commonly diagnosed unintentionally by histopathological examination of ovarian tissue after an ovariectomy for solid ovarian mass.

Ovarian leiomyoma probably arises from smooth muscle cell in the ovarian hilar blood vessels, but other possible origin are cells in ovarian ligaments, smooth muscle cells or multipotential cells in the ovarian stroma, undifferentiated germ cells and cortical smooth muscle metaplasia.3 Ovarian leiomyoma concomitantly seen with uterine leiomyoma (78%), that suggest an identical hormonal stimulation.4

In some instance, MRI can help to determine the preoperative diagnosis, but it may be impossible to delaminate the precise origin of large tumor (uterus subserous versus ovary) on MRI. The tumor appeared as well circumscribed low signal intensity mass on T1 weighted imaging, with mixed signal intensity on T2 weighted imaging. Areas of high intensity signal on T2 weighted images corresponded to degeneration f leiomyoma.5

CASE REPORT

A 29 year old p1L1 women was referred to our hospital for lower abdominal pain and palpable mass at lower abdominal region for 6 months. On examination she was anemic, abdomen shows mass with smooth margin and surface, firm consistency of size 5*5 cm in left iliac fossa. On vaginal examination uterus was normal size and anteverted, left fornical mass firm to hard consistency of 7*6cm size freely mobile with minimal tenderness on bimanual examination. a groove felt between mass and uterus. On CT SCAN a well-defined heterogeneously enhancing mass arising from left anterior wall of uterus, left ovary could not separately
visualized concluding possibility of pedunculated myoma uterus or solid left ovarian mass; she undergone laprotomy.

**Figure 1:** Cut section of ovarian fibroid showing greyish white whorled areas.

**Figure 2:** Intra-operative picture of ovarian mass.

**Figure 3:** Magnified microscopic view of ovarian fibroid showing smooth muscle separated by connective tissue.

**Figure 4:** Microscopic view of ovarian fibroid.

During laprotomy on inspection of uterus and adnexes revealed a solid, firm, oval, left sided ovarian tumor with smooth surface approximately 5*5cm. The tumor was distinctly separated from uterus and exhibited no adhesion to or infiltration of surrounding structures. There was no accompanying uterine mass. The right salpinx, ovary was normal on inspection and were left intact (left salpingo-oophrectomy). After surgery histological examination resulted in a diagnosis of ovarian leiomyoma.

**DISCUSSION**

Most ovarian leiomyoma have no symptoms and discover either during routine physical examination, incidently at surgery, or at autopsy. In symptomatic cases that often present in large one, clinical manifestation is abdominal pain, palpable mass, hydronephrosis, elevated CA -125, Meig’s syndrome and polymyositis.6,7

Ovarian leiomyoma often misdiagnoses preoperatively as pedunculated uterine myoma, ovarian fibroma or even ovarian endometrioma.8

Macroscopic and microscopic manifestations of ovarian leiomyoma are very characteristic, but because of its rarity, several other tumors should include in the differential diagnosis. The major differential diagnostic considerations for ovarian leiomyoma is leiomyosarcoma and sex cord stromal tumors, such as fibroma, thecoma, sclerosing stromal tumor.9

Ovarian leiomyomas must be also differentiated from leiomyosarcoma but due to rarity of these tumors histologic features of malignancy have not been well defined. Pathologists have traditionally used criteria that stress the mitotic count, but it is evident that some other criteria, such as cytological atypia and tumor necrosis must be used when considering the possibility of malignancy in smooth muscle uterine tumor.3 Most of the patients undergone a salpingo-oophrectomy or an oophorectomy with or without hysterectomy despite the
young age of affliction, and only minor of patients are submitted to ovarian preserving surgery. 10-12

CONCLUSION

Ovarian leiomyoma is a very rare tumor of unresolved origin. Despite its rarity ovarian leiomyoma should considered in the differential diagnosis of ovarian spindle cell tumors. Appropriate diagnosis in some cases requires additional immunohistochemical analysis for actin and desmin.

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REFERENCES
