Case Report

Accessory Cavitated Uterine Mass (ACUM) - A Rare Cause of Pelvic Pain in Young Women

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Abstract

Background: A rare Mullerian anomaly may be a cause of severe dysmenorrhea and recurrent pelvic pain not responding to medical management in young women.

Materials and Methods: We present a case report of a rare anomaly, accessory cavitated uterine mass (ACUM) / Uterine like mass managed successfully by laparoscopic resection of the uterine mass and suturing of the myometrial defect on uterus at our institute SDMH.

Results: ACUM or Accessory cavitated uterine mass earlier reported as juvenile cystic adenomyoma represent a rare, newly recognized Mullerian anomaly observed in young females suffering from severe dysmenorrhea and recurrent pelvic pain. The diagnosis is made when an isolated extra-cavitated uterine mass is seen in an otherwise normal appearing uterus with unremarkable endometrial lumen and adnexal structures. Laparoscopic excision of the mass remains the mainstay of treatment.

Conclusion: Awareness about the entity, early diagnosis and laparoscopic excision of the mass brings instant relief to the young woman.

Keywords
Pelvic Pain, Dysmenorrhea, Accessory and Cavitated Uterine Mass; Juvenile Cystic Adenomyoma; Mullerian Anomalies; Uterus-like Mass.

Introduction

Accessory cavitated uterine mass (ACUM) is a rare developmental mullerian anomaly that is characterized by presence of accessory cavity in uterus in an otherwise normal uterus seen in young female presenting with chronic pelvic pain and dysmenorrhea [1-6].

Case report

We report a case of 17-year-old girl who presented to the department of Obstetrics & Gynecology with complaints of recurrent abdominal pain in left iliac fossa since two years. She took medical treatment elsewhere with non-steroidal anti-inflammatory drugs (NSAIDS) and oral contraceptive pills (OCP) for the last few months but did not respond. She attained menarche at the age of 14 years thereafter her menstrual cycles were regular with normal flow. The general physical examination was unremarkable. The history revealed that she underwent ultrasonography examination multiple times before presenting to our institute. The previous ultrasonography reports document different pathologies like ovarian torsion, degenerated myoma and endometrosis. We decided to get a MRI scan done that showed normal uterus with well defined, non communicating cavitated mass measuring 3 x 4 cm along left anterior uterine wall just below the insertion of round ligament was seen. The cavity lining was hyper intense on T2 with hemorrhagic contents that differentiated the
mass from fibroid, adenomyoma as they do not show up hyper intense T2 as they lack endothelial lining (Figure 1). The junctional zone of the cavity was thickened (13mm) with indistinct endomyometrial interface. The uterine cavity per se was normal with normal coruna, ovaries and fallopian tubes without any evidence of endometrotic deposits.

A provisional diagnosis of ACUM was made and the patient was taken for diagnostic Laparoscopy. For diagnostic laparoscopy after creating pneumoperitoneum, 10 mm supra umbilical port was inserted using Verre’s needle along with two 5 mm left accessory side ports. The laparoscopy showed a broad based uterus having mass attached to left anterior uterine wall below the attachment of round ligament. Both ovaries and fallopian tubes appeared normal without any evidence of endometriosis (Figure 2). The patient being an unmarried female hence hysterosalpingography (HCG) was not done. Laparoscopic resection of ACUM was planned immediately after diagnostic laparoscopy confirmed the previous MRI findings. Vasopressin (20 units in 200 ml normal saline) was injected with a needle along the ACUM – uterine serosal interphase in the adjacent myometrium to achieve homeostasis (Figure 3).

ACUM seros was incised using scissors with an incision over anterior wall of mass with subsequent drainage of 10 ml of chocolate colored fluid. We performed Enucleation in a circumferential manner along the ACUM and uterine myometrial interphase using scissors (Figure 4). After resection homeostasis was achieved and uterine defect was closed in 2 layers using 1-0 vicryl (Figure 5). The resected tissue was sent for detailed histopathological examination. The microscopy was indicative of uterine mass comprising of endometrial glands and stroma (Figure 6).

**Discussion**

The first case of such uterine mass was reported by Cozzutto in 1981 [6]. But since thereafter there has been increased awareness among clinicians and it is been diagnosed more frequently. ACUM is a rare congenital mullerian anomaly commonly seen in young nulliparous women presenting with dysmenorrhoea and infertility. The diagnosis of this anomaly is challenging as it confused with several gynaecological disorders including non...
communicating rudimentary uterine horn, true cavitated adenomyoma and generating fibroid. It is characterized by the presence of normal uterus with duplication and persistence of ductal mullerian tissue at the insertion of the round ligament due dysfunction of gubernaculums [5]. It is important that the gynecologist should have high index of suspicion for ACUM where young female complaints of dysmenorrhea and or chronic pelvic pain. The mass lesion in our case was highly suggestive of ACUM considering younger age of presentation with classical complaints of dysmenorrhea and or chronic pelvic pain. The mass lesion was excised and sent for complete histopathologic examination. Grossly, it consisted of multiple grey white soft tissue pieces measuring in total 3.5 X 2.5 X 2 cm with microscopic examination revealing endometrial glands and stroma. The lining of accessory cavity is by endometrial epithelium with glands and stroma. It is characterized by the presence of normal uterus with duplication and persistence of ductal mullerian tissue at the insertion of the round ligament due dysfunction of gubernaculums [5]. It is important that the gynecologist should have high index of suspicion for ACUM where young female complaints of dysmenorrhea and or chronic pelvic pain. The mass lesion in our case was highly suggestive of ACUM considering younger age of presentation with classical complaints of dysmenorrhea, pelvic pain and latter MRI findings showing normal uterus and absence of endometriotic features [3]. ACUM was previously known as Juvenile cystic adenomyosis which is a rare variant of adenomyosis having large hemorrhagic cyst with an ectopic endometrial glands occurring in a young female with dysmenorrhea and pelvic pain that is resistant to medical therapy. There are closely related pathologies that present with dysmenorrhea. Juvenile cystic adenomyoma is a solitary myometrial cyst generally more than 1 cm that is independent of uterus and is lined by hypertrophied endometrium, it presents with dysmenorrhea in young female of less than 30 years of age as a variant of adenomyosis [6]. Complete EN bloc resection with sufficient margin is to be done as treatment. In our case there was absence of adenomyotic lesion and complete resection of mass along with endometriotic lining brought symptomatic relief to the patient [9]. Laparoscopic surgery is the best treatment and anterior approach is preferred over posterior one as it carries lower risk of hemorrhage from ascending branch of uterine artery [10, 11]. The initial investigation is done for evaluation of pelvic pain is ultrasonography USG. On USG it is seen as solid cystic mass with the uterus with clear separate visualization from ovaries and fallopian tubes. HSG has limited value in making concrete diagnosis and is only helpful to rule out mullerian anomalies. MRI is investigation of choice which is preferred over HSG specially in young unmarried female [3]. MRI enables visualization of pelvic anatomy, the cavitated mass along with the presence of hemorrhage contains uterus. MRI also adds in differentiating ACUM from adenomyosis and pelvic endometriosis. After resection the excised mass was sent for complete histopathologic examination. Grossly, it consisted of multiple grey white soft tissue pieces measuring in total 3.5 X 2.5 X 2 cm with microscopic examination revealing endometrial glands and stroma. Table 1 shows a helpful mnemonic for quick recap of this congenital anomaly characterized by three – clinical features, pathology, investigations, treatment options and differential diagnosis- three's anomaly.

**Acien et al described ACUM with following diagnostic criteria:**

- An isolated accessory cavitated mass usually located under round ligament
- Normal uterus, fallopian tubes, and ovaries

**Conclusion**

ACUM or uterus in uterus is a very rare congenital disease that may present with incapacitating pain. For practicing gynecologists it is prudent to keep ACUM in differential diagnosis in young girl with severe dysmenorrhea that is refractory to medical treatment. The extreme pain adds to
the severe mental stress therefore awareness at the level of gynecologist and radiologist is essential as accurate diagnosis and surgical management gives complete cure and improves immensely the quality of life.

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No

**References**


