Hernia uterine inguinale with endometriosis in adult Mayer Rokitansky Kuster Hauser syndrome

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Abstract
A young 25 year old woman presented with primary infertility and was found to have a big left inguinal mass with a blind vagina. Based on the findings of magnetic resonance imaging, a history of probable primary amenorrhea and absence of cervix on clinical examination she was diagnosed as MRKH with inguinal herniation of uterus and adnexae. Operative findings showed bilateral grossly distorted inguinal masses with evidence of endometriosis. Postoperative pathology findings demonstrated endometriosis.

Introduction
The Mayer-Rokitansky-Küster-Hauser (MRKH) syndrome is characterized by congenital aplasia of the uterus and the upper part (2/3) of the vagina in women showing normal development of secondary sexual characteristics and a normal 46, XX karyotype. It affects at least 1 out of 4500 women¹. The majority of cases appear to be sporadic, however family cases have also been described. The mode of inheritance seems to be autosomal dominant with an incomplete degree of penetrance and variable expressivity. Type I (isolated) MRKH is less frequent than Type II². Several cases have been reported where the tubes and ovaries have herniated into the inguinal canal. Most cases concern the paediatric population with a congenital hernia sac. Entrapment of adnexae in an indirect inguinal hernia is rare in adult women³.

Case report
A 25 year old lady, born to non-consanguinous parents, came to us with history of primary infertility and secondary amenorrhea, apparently having attained menarche at 16 years, had regular periods until six months back. There was no history of dysmenorrhea. She was married and sexually active since three years. Her secondary sexual characters were well developed. External genitalia were female. Local examination showed a palpable mass in left inguinal region – 8 × 7 cm. The mass was non tender, irreducible, and cystic in consistency (Figure 1). Per speculum showed a blind vaginal pouch 4 cms depth. Cervix was not seen. Per vaginum examination confirmed speculum examination. No uterus or appendages felt in midline. Thyroid profile, prolactin, follicle stimulating hormone, luteinising hormone, testosterone, dihydroepiandrosterone sulphate levels were within normal limits. Cytogenetic report was normal female karyotype of 46 XX. MRI imaging showed a case of Mayer-Rokitansky-Kuster-Hauser syndrome showing wide separation of hypoplastic uterine horns in to bilateral inguinal regions with respective ovary and dilated blood filled fallopian tubes. Left sided mass of 15 × 5.8 cm. Right sided mass measuring 8.8 × 5.7 cm. (Figure 2). Transrectal USG study showed blind vagina measuring about 4.5 cm in length. Renal and skeletal systems were normal. On surgery a right sided endometriotic tubo ovarian mass was found stuck to the right lateral pelvic wall, in the region of the right internal iliac ring. The rudimentary horn was noticed.

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beneath the endometriotic cyst. The tube was distended to form a huge haematosaphinx which had partially prolapsed into the right inguinal canal. The contents were reduced by traction and blunt dissection. The right rudimentary horn was excised with the tubo ovarian mass. On the left side a bigger mass was seen with major portion of the mass extending into left inguinal canal which could not be reduced. Hence a left inguinal incision was taken and a left inguinal hysterectomy was done (Figure 3). Conservative surgery could not be done due to gross distortion of structures. Histopathology of the mass showed rudimentary horn of uterus with adenomyosis, chocolate cyst of ovary compatible with endometriosis and dilated fallopian tube showing hemosiderin laden macrophage, chronic inflammatory cells, congested blood vessels and fibroblast, giant cells. No cervical tissue was seen.

Discussion

Inguinal hernia containing uterus and endometriosis is exceedingly rare. Most inguinal endometriosis is located at an extra pelvic site near the round ligament. One such reported case of a patient with inguinal hernia containing rudimentary uterine horn and endometriosis was unilateral5. A handful of other cases show a unilateral herniation of uterus, tubes and ovaries without endometriosis56. Several cases of ovarian endometriosis in MRKH without uterine hernia have been reported58. Our patient was found to have bilateral herniation and endometriosis. Endometriosis in a patient with MRKH syndrome with a small rudimentary uterine horns and functional endometrium can be assumed to have developed by retrograde menstruation theory. Such patients usually present with cyclical pain during menstruation as reported in the one case5. Surprisingly our patient did not have any pain nor did she acknowledge the evident primary amenorrhoea. Psychosocial elements should be considered when dealing with
such patients. Hernia uterine inguinal has been reported most commonly in the literature as both persistent müllerian duct syndrome and male pseudohermaphroditism. It is most often seen in a phenotypically normal male infant having both testes and uterine tissue present. Few cases have been documented to occur in the female sex or the adult patient. This subset may be at higher risk for hernia uterine inguinale, and if presenting with complaints of pain or inguinal mass, it should likewise be considered in the differential diagnosis. Abdominal and pelvic imaging is useful in the diagnosis of this condition because it may aid in identifying patients with coexisting müllerian malformations9.

REFERENCES