Incidental Finding of Leiomyoma in Mayer-Rokitansky-Kuster-Hauser Syndrome

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Abstract
Mayer-Rokitansky-Kuster-Hauser (MRKH) syndrome is a sexual developmental disorder. In this disorder, there is a congenital absence of the uterus and vagina with normal external genitalia. The etiology is not well understood. Variations of this condition exist that may include congenital abnormalities and psychological problems. In this article, we discuss the case of a 47-year-old African American female who presented with acute renal failure, solitary right kidney, and a pelvic mass extending from the pelvis to the right hypochondrium determined to be a fibroid. The patient was managed by a multidisciplinary team, dialyzed, and planned for removal of the mass. While understanding the low probability of having fibroids without a uterus, fibroids should not be excluded from such patients. It is also important to consider the emotional and psychological well-being of such patients.

Keywords
Mayer-Rokitansky-Kuster-Hauser syndrome, MRKH, leiomyoma, Mullerian duct

Introduction
Mayer-Rokitansky-Kuster-Hauser (MRKH) syndrome is associated with an absence or incomplete development of the Mullerian duct.1 Three different types of this syndrome may exist: (1) the uterus and upper vagina are absent; however, fallopian tubes and normal ovaries exist; (2) ovarian or kidney malformation and no aplasia or hypoplasia of the uterus and vagina; (3) kidney dysfunction in association with other congenital anomalies (skeletal, cardiac, and auditory).2 Familial clustering of this syndrome in conjunction with congenital anomalies may suggest a genetic link. While hormone levels may be normal in these patients, there may be hormone deregulation in which some patients have hyperandrogenemia and hyperprolactinemia. Luteinizing hormone (LH) and follicle-stimulating hormone (FSH) are normal in MRKH syndrome, which may indicate appropriate ovarian function. Testosterone levels are in the normal female range.

The incidence is 1 in 5000 female births.3 As a result of hormonal dysregulation, secondary sex characteristics may be affected with resultant amenorrhea during the onset of puberty. This condition is the second most common cause of primary amenorrhea after ovarian failure in reproductive aged women.4 Psychological issues include low self-esteem, emotional distress, depression, rejection, among others.5-7 Very few cases of MRKH have been described in the medical literature.

Case Description
A 47-year-old African American female with HIV, hypertension, hepatitis B, and hepatitis C presented to the emergency room with severe right-sided chest pain, nonproductive cough, and fever for 2 days. She had never had a menstrual period. She smoked one half pack of cigarettes, cocaine, and injected heroin daily for over 10 years. There was cachexia, prolonged expiratory phase on auscultation, and an abdominal mass extending from the pelvis to the right hypochondrium. Pelvic examination revealed a shallow vaginal canal,
and the cervix was not visualized. Chest X-ray showed right basilar infiltrate with a small right pleural effusion. Her blood urea nitrogen (BUN)/creatinine ratio was 40/4.9 (approximately 8). CA-125, α-fetoprotein, human chorionic gonadotropin (hCG), and lactate dehydrogenase (LDH) were all within normal limits. Colonoscopy was negative. On admission, the patient received azithromycin and felt better though found to have worsening renal function with BUN/creatinine persistently trending upward—BUN/creatinine at 63/5.9 and was started on dialysis. Computed tomography (CT) scan of abdomen and pelvis was done. Cystoscopy revealed obstruction of the right ureter. Right nephrostomy tube was placed to relieve obstruction.

CT scan showed well-circumscribed complex pelvic mass measuring 19 cm by 11.7 cm by 9.6 cm, and a solitary right kidney (see Figure 1). Pelvis mass biopsy revealed a well-differentiated smooth muscle tumor with no atypia or mitotic activity (see Figure 2). Tumor cells were positive for desmin and estrogen receptors, favoring smooth muscle origin. Pelvic ultrasound was consistent with a diagnosis of MRKH, showing a blind vagina and little to no uterus, with both ovaries appearing normal. MRKH syndrome was a top differential considering her solitary kidney and primary amenorrhea and excluded by pelvic scan and hormonal profile were the following: (LH and FSH) primary amenorrhea, androgen insensitivity syndrome (shallow vaginal canal and the cervix not visualized), cervical dysgenesis (vaginal examination), and gonadal dysgenesis (Turner’s syndrome-no stigmata). No skeletal abnormalities were seen by magnetic resonance imaging (MRI). Attempts to replace nephrostomy tube with intra-ureteral stent were unsuccessful due to persistent compression by pelvic mass. She was discharged with stent in situ and scheduled for dialysis 3 times per week and for follow-up with gynecology within 1 week for removal of fibroid tumor for which she declined.

**Discussion**

In a patient with MRKH syndrome, the proximal two thirds of the vagina do not develop and patients have some of the VACTERL-associated conditions that include but are not limited to the vertebra, heart, urogenital, and ear anomalies. Secondary sexual characteristics are present with a 46 XX karyotype. In order to make a definitive diagnosis, a clinical examination followed by pelvic ultrasonogram and MRI, and karyotype are required. For many patients, the initial presentation is normal female sexual characteristics with primary amenorrhea. According to medical literature, there are many causes of amenorrhea and MRKH syndrome is the second most common cause.

This syndrome has gone through multiple revisions in its description to capture the entire picture between 1829 and 1961. In 1829, Mayer described various vaginal anomalies not limited to duplications because of abnormal development of the Mullerian ducts. In 1838, Rokitansky described uterine and vaginal agenesis. Kuster in 1910 recognized renal abnormalities (renal ectopy or agenesis) and skeletal abnormalities. Hauser made the distinction between MRKH and testicular feminization in 1961. It may either be sporadic, familial, or genetic in etiology. Transmission is autosomal dominant, and multiple patients with this genetic condition may have variable presentation. A link between a loss-of-function mutation in the WNT4 gene and MRKH has been described in the medical literature.

We present a case of Mullerian agenesis in a patient with a biopsy-confirmed large fibroid tumor, from the vestigial Mullerian remnant that grew large enough to obstruct her only right kidney resulting in dialysis dependency.
Our patient who had worsening renal function, solitary kidney, amenorrhea, and a pelvic mass was incidentally discovered to have MRKH with leiomyoma at index presentation with no documented history of MRKH. Recent cases of MRKH syndrome with fibroids showed that patients varied from ages 20 through 70 years and with no specific country predilection. Twenty-one percent of MRKH patients had a solitary kidney, but unilateral renal anomalies range from 30% to 50%.4,9 The congenital malformations observed in this patient suggested an MRKH-like phenotype.1,11

In a review of the medical literature in 35 patients with MRKH and leiomyoma, the origin of leiomyoma in MRKH patients was the uterine remnant in 65% of patients.4 Other origins include multiple uterine remnants, fibrous myometrial bands, and the broad ligament, round ligament, vascular leiomyoma, and parametrial tissue.4,8 Leiomyomas can originate from the myometrial portion of the uterus. They are known to have fibromuscular tissue comprising smooth muscles (from a study of embryology, Mullerian ducts at the proximal ends have smooth muscle cells but are primarily endodermal in origin) and can grow because of estrogen stimulation; considering that patients with MRKH have ovaries with normal hormone levels, fibroids in these patients undergo the same patterns like in normal patients. In addition, it has been suggested that estrogen receptor sensitivity or concentration in the remnant tissue may be potentially responsible. Surgical or nonsurgical treatment for MRKH is available, but this is dependent on individual needs, patient motivation, and available options.13 There are 2 main types of procedures: (1) A neovagina is created for patients who have signified psychological readiness for sexual activity.11 The physiological defects are both medically and surgically treatable, permitting sexual function.13 With assisted reproductive techniques and surrogacy, MRKH syndrome patients can reproduce.14 (2) An already identified and prepared canal lined with bowel as mucous membrane is utilized for vaginal replacement.15 Adherence to a home dilation schedule and recommending consistent condom use to prevent infections such as HIV and human papilloma virus are key. Regular examinations should be performed to assess for vaginal stenosis.15

The patient’s polysubstance abuse made her unstable in the hospital setting and complicated any inpatient plans to treat her and this reflected in her subsequent loss to follow-up. There is a growing need to explore the emotional and psychological well-being of MRKH patients because of their unique experiences.5,7 because they feel rejection, depression, hindrance of independence in younger women, loss of self-esteem, anxiety, loss of identity, and feeling other than. Many patients receive a diagnosis of MRKH because of primary amenorrhea in adolescence. It is appropriate to receive psychological intervention during the transition from adolescence to adulthood; however, our patient did not receive this.5,7 Management should be multidisciplinary and should be tailored to specific needs for specific patients.

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Ethics Approval
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Informed Consent
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