Herlyn-Werner-Wunderlich syndrome: a rare cause of acute urinary retention in an adolescent girl

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DESCRIPTION

A 14-year-old girl presented to emergency triage with acute retention of urine. There was history of cyclic pelvic pain for the past 6 months, for which she had been prescribed non-steroidal anti-inflammatory drugs by a private practitioner. There was no history of ureteric colic or burning during micturition. Gynaecological history indicated menarche at 12 years followed by regular menses. There was no history of abnormal vaginal bleeding. Ultrasound of the abdomen revealed findings suggestive of bicornuate uterus with haematocolpos. Subsequently, MRI was requisitioned, revealing two widely divergent, symmetrical uterine corpii, partially fused at the cervix without any communication between their endometrial cavities (figure 1A). There was a thick hypointense left hemivaginal septum with a large ipsilateral haematocolpos appearing hyperintense on T1-weighted and hypointense on T2-weighted images, findings consistent with intracellular methaemoglobin stage of bleed. A similar intensity dilated tortuous...
tubular structure was seen extending from left uterine cornu up to the left ovary, findings consistent with haematosalpinx. The right uterine cervix was seen normally opening into the patent right hemivagina (figure 1B, C). In addition, left renal agenesis with compensatory hypertrophy of the right kidney was noted (figure 1D). Bilateral ovaries were normally visualised with a small haemorrhagic cyst seen in the left ovary. A diagnosis of Herlyn-Werner-Wunderlich syndrome (HWWS) was performed and the patient underwent laparoscopic vaginal septectomy, where 500 mL of chocolate coloured fluid was drained. The patient became symptom free after the procedure.

HWWS is a rare, combined Mullerian and Mesonephric duct anomaly characterised by the triad of uterus didelphys, obstructed hemivagina and ipsilateral renal agenesis.1 Uterus didelphys represents failure of fusion of the paired Mullerian ducts. Obstructed hemivagina results from a transverse vaginal septum, which represents vertical fusion defect consequent to non-resorption of tissue between the vaginal plate that originates from the urogenital sinus and the caudal aspect of the paired Mullerian ducts. 2 The renal agenesis on the same side of the obstructed hemivagina reflects the close embryological association of Wolffian duct in development of Mullerian ducts, whereby developmental anomaly of one Wolffian duct may cause unilateral renal agenesis associated with obstructed hemivagina. Unlike our case, which shows a left-sided obstructed system, a unilateral renal agenesis associated with obstructed hemivagina.

The reported incidence of uterine didelphys with renal agenesis in the general population is 0.1–3.8%. The most common presenting symptoms of HWWS are cyclical dysmenorrhea at the onset of puberty, associated with vaginal or pelvic mass. Rarely, it may have delayed presentations of primary infertility, ischiorectal swelling, pyometra or urinary obstruction.1 2

In women, the urethra has a small length, straight course and large diameter and urinary retention is therefore very rare. Fewer than 10 cases of acute urinary retention have been ascribed to haematocolpos.7 Acute urinary retention in the setting of haematocolpos can be explained by its mass effect on the urethra causing its stretching or angulation and leading to its obstruction. Another hypothesis is that haematocolpos may cause diminished extruding capacity of the extruder muscle or an increased contractility of the urethral sphincter through the irritation of the sacral plexus base.4

MRI is the investigation of choice for suspected Mullerian duct anomalies, as it exquisitely defines the internal and external uterine anatomy, and accurately assesses the complex anomalies, thereby helping diagnose secondary conditions such as fibroids, endometriosis and hydrosalpinx and providing a surgical road map for further management.

The ideal treatment for HWWS is laparoscopic vaginal septum resection and/or septoplasty, but menstrual suppression by oral contraceptive pills or gonadotropin-releasing hormone analogues is advised if immediate surgery is not an option.8

Owing to its rarity, HWWS is usually not included in the differentials for acute urinary retention in a young woman, but its early recognition and management can prevent further complications such as endometriosis, pelvic adhesions and infertility. Thus, we would re-emphasise the need for sensitisation of healthcare professionals, especially in emergency units, to this rare entity, HWWS.

Learning points

► Herlyn-Werner-Wunderlich syndrome (HWWS) is a rare, combined Mullerian and Mesonephric duct anomaly characterised by the triad of uterus didelphys, obstructed hemivagina and ipsilateral renal agenesis.

► A high suspicion index is warranted for early recognition and prompt treatment in order to prevent future serious complications such as endometriosis, pelvic adhesions and infertility.

► In adolescent girls, HWWS can present with acute urinary retention and hence it should be considered as a differential in such a clinical scenario.

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