Herlyn-Werner-Wunderlich syndrome with a partially obstructed hemivagina

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Uterine didelphys with obstructed hemivagina and ipsilateral renal agenesis is a rare congenital anomaly of the Müllerian duct system referred to as Herlyn-Werner-Wunderlich syndrome. Because of its rare occurrence, a high level of suspicion is often required for diagnosis. Clinically, these patients usually present after menarche with pelvic pain, dysmenorrhea, and a palpable pelvic mass. We present a case of a 31-year-old female patient with infertility. Imaging findings were consistent with Herlyn-Werner-Wunderlich syndrome, with a congenital defect in the longitudinal vaginal septum resulting in partial spontaneous decompression of right-sided hematocolpos.

**Case report**

A 31-year-old female presented with 18 months of primary infertility. Her menstrual cycles were regular. She had no history of dysmenorrhea, pelvic mass, or pelvic pain. A bimanual pelvic examination yielded a normal-size uterus that was mobile and nontender, without palpable pelvic masses. A speculum examination demonstrated a nulliparous cervix without lesions.

The patient reported that a prior evaluation elsewhere had indicated a septate uterus or uterine didelphys. Magnetic resonance imaging (MRI) of the pelvis (to evaluate the possible Müllerian anomaly) was performed on a 1.5-Tesla MR scanner (Achieva, Philips Medical Systems). An intravenous contrast agent was not considered necessary. Images were acquired on multiple planes with oblique axial T1 (TR 608 ms; TE 10 ms), oblique coronal T2 (TR 3733 ms; TE 120 ms), oblique axial T2 (TR 3172 ms; TE 120 ms), sagittal T2 (TR 8276 ms; TE 120 ms), and coronal T2 SPAIR (TR 761 ms; TE 120 ms) sequences.

**Figure 1.** 31-year-old female with Herlyn-Werner-Wunderlich syndrome. T2-weighted axial MR image through the pelvis demonstrated two uterine horns (black arrows) and two individual cervices (white arrows). The uterine horns were almost of normal size, completely separate, and widely splayed.

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MR images showed two uterine horns that were completely separate and widely splayed. They appeared fully developed and almost normal in size, with preserved endometrial, myometrial, and junctional zones. Two individual cervices were in continuity with the uterine horns (Fig. 1). A longitudinal vaginal septum divided the upper vagina into two cavities, with one cervix entering each hemivagina. A defect in the proximal end of the vaginal septum allowed direct communication between the two hemivaginas (Fig. 2). The right hemivagina was moderately distended and contained blood products that were T2-hyperintense and mildly T1-hyperintense, consistent with hematocolpos. The right hemivagina terminated blindly approximately 4 cm cranial to the introitus. The left hemivagina was decompressed (Fig. 2). A single lower vaginal cavity likely communicated with the left hemivagina, the left cervix, and the left uterine horn. Coronal images through the midabdomen demonstrated an empty right renal fossa and a normal left kidney (Fig. 3). These imaging features were diagnostic of HWW syndrome with a congenital defect in the longitudinal vaginal septum, resulting in partial spontaneous decompression of the right hemivagina.

Discussion

Herlyn-Werner-Wunderlich (HWW) syndrome is a congenital anomaly of the Müllerian and Wolffian ducts (1) characterized by a triad of uterus didelphys, obstructed hemivagina, and ipsilateral renal agenesis (2-5). The diagnosis is usually made soon after menarche due to symptoms related to the obstructed hemivagina, such as pelvic pain, dysmenorrhea, and palpable mass from associated hematocolpos or hematometra.

An obstructed hemivagina with a didelphys uterus is the most common type of obstructing malformation of the uterus and vagina (6). The longitudinal vaginal septum is due to incomplete disappearance of the partition between the fused Müllerian ducts (7). As an obstructing vaginal malformation, this condition precludes the outflow of menstruation, resulting in hematocolpos, possible hematometra, and hematosalpinx. The diagnosis is therefore usually made soon after menarche due to symptoms related to the obstructed hemivagina, such as pelvic pain, dysmenorrhea, and a palpable pelvic mass. Long-term complications of HWW syndrome include endometriosis from retrograde menstruation, and obstetric complications such as recurrent pregnancy loss, preterm labor, abnormal fetal presentation, and prematurity due to uterine anomalies (8-10).

Our patient presented with primary infertility and had a congenital defect in the longitudinal vaginal septum, resulting in partial spontaneous decompression of the rightxed hemivagina. As illustrated by this case report, patients with HWW syndrome may have a defect in the longitudinal vaginal septum that allows spontaneous decompression of the obstructed hemivagina into the nonobstructed hemivagina. Such patients are likely to present with fewer obstructive symptoms than in those with an intact longitudinal vaginal septum and complete obstruction of the hemivagina. Although some amount of hematocolpos and/or hematometra may be present at diagnosis, we as...
sume that the volume may not be sufficient to cause ob-
struc-tive symptoms. This would explain the delay in presen-
tation as in the case presented. We also postulate that pa-
patients with a defect in the longitudinal vaginal septum may
be protected from retrograde menstruation and subsequent
development of pelvic endometriosis, since the hemivagina
is partially decompressed via the defect.

To our knowledge, this is the only reported case where a
defect in the septum has resulted in decompression of the
obstructed vagina. MRI has been shown to be the gold
standard in imaging of Müllerian anomalies (1), and we
consider it the imaging modality of choice for illustration of
defects in the longitudinal vaginal septum.

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