Partial Vaginal Agenesis with Transverse Vaginal Septum

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ABSTRACT
Partial vaginal agenesis with transverse vaginal septum is rare malformation of female genital system. The reported incidence of vaginal atresia is 1:4000 to 5000 live female birth. The incidence of septum in upper one-third is 46%, middle 35%, lower 19%. Transverse vaginal septum occurs due to defect in vertical fusion of Mullerian ducts during embryogenesis. Vaginal atresia occurs due to failure of canalization of sinovaginal bulbs.

We are presenting such a rare case of partial vaginal agenesis with transverse vaginal septum.

The aim of this presentation is to make the consideration of transverse vaginal septum and vaginal atresia in young girls with cryptomenorrhea and to emphasize on the meticulous evaluation and diagnosis with counselling of a minor along with parents of future reproductive outcome along with surgical difficulties anticipation and difficulties post vangioplasty.

CASE REPORT
A patient aged 17 years presents with complaints of cyclical pain abdomen. Pain used to be present once in a month cyclically. No family history of similar complaints. Patient had well developed secondary sexual characters. Height was 146 cm and no features of Turner’s syndrome were present and breast development was Tanner stage 4, pubic hair and axillary hair were present, external genitalia was normal. Hymenal opening was not seen and dimple was present in hymenal area. On abdominal examination, a suprapubic bulge was present and it was tender.

Her hormonal analysis was within normal limits, echocardiography (ECHO) normal, X-ray spine normal. Ultrasonography report showed uterus 123 × 41 × 74 mm, evidence of hetero-echoic collection in the cervix and the uterine cavity. Collection of around 300 ml in the vagina. Thick septum in lower part of vagina. Lower 1/3 part of vagina was not made out. No renal anomalies were found in the ultrasound (Fig. 1). Magnetic resonance imaging (MRI) was not done as patient was not affordable. Karyotyping was 46 XX.

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INTRODUCTION
Partial vaginal agenesis with transverse vaginal septum is rare malformation of female genital system. The reported incidence of vaginal atresia is 1:4000 to 5000 live female birth. The incidence of septum in upper one-third is 46%, middle 35%, lower 19%. Transverse vaginal septum occurs due to defect in vertical fusion of Mullerian ducts during embryogenesis. Vaginal atresia occurs due to failure of canalization of sinovaginal bulbs.

We are presenting such a rare case of partial vaginal agenesis with transverse vaginal septum.

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Fig. 1: Ultrasound picture of partial agenesis of vagina and distended upper vagina and uterus with collection
Clinical diagnosis of partial vaginal agenesis with transverse vaginal septum was made. Patient and attendees were counselled regarding necessity of surgery, reproductive outcome and coital functions and need of further follow-ups. Case discussed with surgeon and surgical gastroenterologist. In collaboration with them, patient was planned for reconstruction of vagina and incision of septum and drainage of hematocolpos and hematometra. A space was created between bladder and rectum, and space was made till the level of septum (Fig. 2). The septum was incised under ultrasound guidance and approximately 500 ml of collected blood, was drained. As the septum was distended with blood, it was stretched and there was sufficient tissue and septal edges were pulled through and posterior wall was sutured to introitus and anteriorly pulled out and stitched at the level of vaginal introitus (Fig. 3). Intracervical Foley’s inserted for the drainage of hematometra and mold was inserted into the vagina. Postoperatively mold was changed everyday. Postoperatively on day 10, a small speculum was easily inserted and the patient was advised on need of dilatation with mold repeatedly and discharged (Fig. 4). She had menstrual flow for 4 days in next cycle and ultrasound done after menstrual cycle showed patent tract and no collection in vagina and uterus.

**DISCUSSION**

Development of female genital tract is a complex process; it is dependent on the events involving cellular differentiation, migration, fusion, canalization. Between the sixth and eighth weeks of fetal development, the caudal aspects of the bilateral Mullerian ducts fuse at the midline. A single midline tubular structure forms as a consequence of cell proliferation (i.e. uterovaginal canal). This midline structure extends to the Mullerian tubercles, where it encounters the urogenital sinus. Bilateral endodermal invaginations (i.e. sinovaginal bulbs) form as the Mullerian tubercles regress. Canalization of the uterovaginal canal is believed to occur from the caudal to the cephalic aspect, with an epithelial lining derived from the urogenital sinus. Failures at the vaginal plate level explain transverse vaginal septum and its variant.

Partial vaginal atresia and transverse vaginal septum occur due to defect in above process. Usual age group of presentation of such anomalies is 19 to 38 years in a girl with well developed secondary sexual character. Transverse vaginal septum follows an autosomal recessive mode of inheritance. It is associated with genitourinary, gastrointestinal, musculoskeletal and cardiac anomalies. Genitourinary and gastrointestinal anomalies include imperforate anus, malformation of gut, ectopic ureter with hypoplastic kidney, hydronephrosis, musculoskeletal include malformation of lumbar spine and sacral hypoplasia. Cardiac anomalies include coarctation of aorta and ASD.

Usually, they present during adolescent age group with cyclical pain abdomen, suprapubic bulging, very
rarely urinary retention, outflow incontinence, hydronephrosis. If septum is perforated without vaginal atresia, they may be asymptomatic or with or without dysmenorrhea, infertility, menstrual irregularities, dyspareunia, soft-tissue dystocia. Very rarely if may present at neonatal period with mass per abdomen and pain abdomen. The main treatment modality is surgical. Outcome needs to be considered in such cases is relief of symptoms and sexual function and pregnancy.

Imaging plays an indispensible role in diagnosis of these anomalies. Transverse vaginal septum is pink, thick and does not allow transillumination on clinical examination. On ultrasonography, hourglass distension of upper segment uterus and vagina is made out and sometimes even hematosalphinx can be made out. Magnetic resonance imaging is definitive method.

In our case, we have created the space between the bladder and rectum, septum was incised, and both edges are pulled down and sutured to introitus. Patients own tissue was used so that chances of graft were avoided. Further need of dilatation was emphasized. Early marriage and child bearing was advised to the parents, as they were hematometra and hematosalphinx and chances of endometriosis was explained. Reproductive outcome depends on location of septum retrograde spillage leading to hematometra, hematosalphinx and endometriosis. Coital function depends on the length of vagina reconstructed. Complications of vaginoplasty include injury to the bowel, bladder, infection, blood loss, fistula formation, restenosis, sexual problems and problems with lubrication.

In our study, we want to emphasize that as much as possible we should not resect the septum, rather we should use it for the reconstruction of vagina. Advantages of using the stretched septum for the reconstruction of vagina are since it is an epithelized tract chances of restenosis are very less and graft will take up very nicely and no use of external tissue so there is less chances of infection.

REFERENCES