Pregnancy in a woman with untreated bladder exstrophy: a case report

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Abstract

Objective: To report the management of urinary tract obstruction and infection in a pregnant woman with unrepaired bladder exstrophy.

Case Report: A 27-year-old pregnant woman with unrepaired bladder exstrophy was referred to our hospital with a complaint of bilateral flank pain in the second trimester. After two-dimensional abdominal ultrasound, magnetic resonance imaging and a urine analysis, she was diagnosed with an upper urinary tract infection due to ureteral obstruction secondary to unrepaired congenital bladder exstrophy and an intrauterine pregnancy. J-tube insertion was performed after locating the ureteral orifices and antibiotics were administered. Symptoms rapidly resolved. She delivered a normal male infant by caesarean section at 34 weeks of gestation.

Conclusion: Standard urological management of the ureteral obstruction in pregnancy was successful in this extreme case of unrepaired bladder exstrophy associated with an intrauterine pregnancy. The perinatal outcome was good.

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1. Introduction

Bladder exstrophy (BE) is a rare congenital anomaly of the genitourinary tract that is in the spectrum of the exstrophy–epispadias complex. The embryological mechanism is unknown, though it is thought to be a ventral fusion error due to failed reinforcement of the cloacal membrane by the underlying mesoderm [1]. Its most visible consequence is protrusion of the dorsal mucosal surface of the urinary bladder (trigone and ureteral orifices exposed) through a defect in the abdominal wall. The clinical presentation frequently will include other ventral fusion defects, including of the pubic symphysis, pelvic musculature and genitalia [2,3]. The function of the urinary and reproductive systems is often affected. The benefits of exstrophy repair include moving the bladder into the abdomen (with a reconstructed ventral wall), developing control of urination, reducing the risk of infection, reducing the risk of urinary obstruction, improving physical appearance (self-esteem) and reducing the risk of infertility [2]. Several centres have collectively reported more than 130 corrected BE patients, experiencing more than 100 pregnancies. These data include more than 55 caesarean sections, with 4 stillbirths and 1 neonatal death [4–6]. However, rarely has pregnancy been reported in non-reconstructed BE patients [7,8].

Here, we present the case of a pregnancy in a woman with untreated BE, who in addition developed urinary tract obstruction and upper tract infection in the second trimester.

2. Case Report

A 27-year-old pregnant woman (gravida 1, para 0) with unrepaired bladder exstrophy was referred to our hospital with a chief complaint of bilateral flank pain in the second trimester. The patient was diagnosed with congenital BE but reconstructive surgery was refused by her family. Her regular menstrual cycles had started at the age of 13 years. She had no medical complaint in relation to sexual intercourse and conceived normally. She had not received any antenatal care before this hospital visit. Her medical history and family history were unremarkable.

Physical examination revealed her vital signs were stable. Heart, lung and neurological examinations were unremarkable. Her body mass index (BMI) was 22.6 kg/m². The abdomen was enlarged secondary to a 21 1/2-week pregnancy. The pubic symphysis was not fused, resulting in a 14 cm separation. The exposed mucosa of the posterior bladder wall occupied the lower central abdomen with absence of an umbilicus, urinary meatus, anterior abdominal wall and anterior bladder wall. The mucosal edges of the bladder were fused with the skin. Paroxysmal urine egressed from the exposed ureteral orifices. The vaginal introitus was small and presented as a line approximately only 2.5 cm in length. Percussion pain was evident over the kidneys bilaterally.
Apgar score was 9, and the 5-minute Apgar score was 10. When the male infant was positioned transverse-breech and delivered weighing 2010 g through a low transverse incision in the uterus. The 1-minute Apgar score was 9, and the 5-minute Apgar score was 10. When she began to experience irregular uterine contractions and light vaginal bleeding, two-dimensional abdominal ultrasound showed a single intrauterine pregnancy (compatible with 33+6 weeks of gestation) in the breech position and second-degree placental maturity. Laboratory data showed mild anaemia. The management strategy included promoting fetal lung maturation with dexamethasone, tocolytic therapy, and antibiotics, along with careful fetal monitoring. After appropriate patient education concerning recurrent obstruction, the patient was scheduled for regular follow-up with colour Doppler ultrasound. Checking the placenta, a partial placental abruption was evident, with a separation area of 4 cm × 5 cm.

The patient was discharged on her 7th post-delivery day with both her and her baby in good condition. She was scheduled for a postpartum evaluation and a discussion of reconstructive therapy for her bladder exstrophy. A post-partum pelvic X-ray was obtained to aid in the planning of the reconstructive surgery. The X-ray revealed that the rami of the pubic bones were chondrified and separated by 14 cm. The two J-tubes were imaged in situ. The left J-tube showed a circuitous trajectory and its upper end was located lower than the position of a normal kidney. This tube remained in a stable position even though it did not reach the location of her left kidney, which was documented by two-dimensional abdominal ultrasound and MRU and to be in a normal position. We believe the J-tube followed this circuitous trajectory as a result of ureterovesical obstruction, which led to pyeloureterectasis and circuitous ureter (Fig. 2).

A repair plan was offered but the patient refused further treatment. Therefore, she was scheduled for radiographic evaluation 6-weeks post-partum to assess the upper urinary tract prior to removal of the J-tubes. After appropriate patient education concerning recurrent obstruction, the patient was scheduled for regular follow-up with colour Doppler ultrasound.

### 3. Discussion

Bladder exstrophy is a rare congenital anomaly that is often accompanied by multiple malformations [1]. Patients who have not undergone reconstructive surgery have more complications. Obstetric complications include infection, prematurity, placental abruption, malpresentations and genital prolapse [1]. Reported urinary complications include: urinary tract infection, urinary obstruction, hydronephrosis, ureteral stricture and ureteric stones [1,2]. In our case all these complications were present except for genital prolapse and ureteric stones. Urinary tract infection is very common due to the defects of abdominal wall and bladder wall. In our case this was diagnosed by symptoms and numerous white blood cells in urinalysis. BE can also lead to stricture of uretero-vesical orifices (as it did in our case) which causes uretero-vesical obstruction followed by pyeloureterectasis and hydronephrosis [1]. This was confirmed by two-dimensional abdominal ultrasound and MRU in our patient. Furthermore, the enlarged gestational uterus aggravated the uretero-vesical obstruction.

Ideally, urinary tract infection and uretero-vesical obstruction in such patients should be monitored by urinalysis, urine culture, colour

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**Fig. 1.** Post-corporeal caesarean section image showing the untreated exstrophic bladder. Operative incision was well healed. Bladder exstrophy was untreated. Umbilicus was absent and inner surface of the posterior wall of the bladder occupied the lower central abdomen. The mucosal edges of the exstrophic bladder were fused with the skin. J-tubes were inserted in both ureteral orifices and connected with urine collecting bags. Vaginal orifice located below exstrophic bladder as line shape and meatus urinarius was absent.

**Fig. 2.** Bone radiography of pelvis. X-ray revealed the rami of the pubic bones were chondrified and separate by 14 cm. Circuitous inserted J-tube image is shown on the left of the image.
Doppler ultrasound exam and even MRU exam. Unfortunately, our patient was non-compliant. Antibiotics were of course important and effective in combination with removal of the obstruction. Extensive surgical intervention should be avoided until the pregnancy has concluded to avoid inducing an abortion or premature labour [9]. In our case the intrarureteral J-tube insertion with the assistance of a ureteroscope was successful, and was the least invasive way to remove the obstruction.

Often, due to the non-closure of the pubic symphysis, vaginal birth is readily achievable in these patients [10]. However, the delivery in our case was more complex. Caesarean section delivery was the only option due to malformations of the birth canal and malpresentation. In addition, the presence of a placental abruption would have added progressive risk. Caesarean section was performed promptly at the onset of labour [3]. It was difficult to decide the position of the incision since the lower central abdomen was occupied by the extrophic bladder. We were concerned that a high incision would make it difficult to deliver the fetal head [7]. We did the caesarean section after choosing a transverse incision of 12 cm at three-horizontal-finger distance above the upper edge of the extrophic bladder, and then made a low transverse incision in the uterus. We assembled a multidisciplinary team so we could respond confidently to any unexpected events.

In conclusion, although BE is rare, pregnancies in women with untreated BE do occur, and are often accompanied by urinary and obstetric complications. Reconstructive surgery should also be offered to improve quality of life.

**Contributors**

All authors have made a substantive contribution to the information or material submitted for publication and take public responsibility.

**Conflict of Interest Statement**

The authors declare that they have no conflict of interest.

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**Consent**

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**Provenance and Peer Review**

This case report was peer reviewed.

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