Incidental Finding of Posterior Urethral Valve during Routine Antenatal Ultrasound: Diagnostic Imaging Case Report in Botswana

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Abstract: Medical ultrasound is a particularly useful imaging method used in the diagnosis of urinary fetal anomalies. One such fetal anomaly is the posterior urethral valve (PUV), which is a congenital malformation of the male urethra occurring early in gestation life. PUV is a common cause of obstructive uropathy in males and fetal death during pregnancy. We report from a medical imaging perspective on a rare and high-risk case of a PUV detected during routine antenatal ultrasound imaging in Botswana. The first scan performed at 16 weeks of gestation age was unremarkable. However, subsequent antenatal ultrasound scans at and after 20 weeks demonstrated bilateral hydronephrosis, distended ureters, and a urinary bladder which gradually increased with the gestation age. There was also a corresponding reduction in amniotic fluid and fetal movements. As a result of these complications, the pregnancy was classified as a “high-risk.” At 30 weeks, spontaneous labour occurred, and a fresh still birth male fetus was delivered.

Keywords: Anhydromnios, Antenatal ultrasound, Hydroureteronephrosis, Posterior urethral valve (PUV).

INTRODUCTION

Medical ultrasound plays an important role in the diagnosis of congenital urinary bladder anomalies during fetal development. One of the fetal anomalies is the posterior urethral valve (PUV). Kilciler et al., [1] define a posterior urethral valve as a congenital malformation of the male urethra occurring early in gestation life. It is the most common cause of congenital bladder outlet obstruction and its responsible for about 17% of pediatric end-stage kidney disease [2, 3]. This condition is present in males with an incidence in the range of 1:5000 to 1:8000 live births [3, 4]. In severe cases of near-complete or complete obstruction during fetal development, death may result due to oligohydramnios, fetal renal failure, and pulmonary hypoplasia [4, 5]. However, those who survive after birth suffer an ongoing bladder dysfunction and progressive kidney injury resulting from early childhood [2, 4, 6, 7]. These complications can be minimized with early diagnosis and appropriate management of the patient in specialised hospitals.

In Botswana, the Ministry of Health (MOH) 2010 guidelines for antenatal care and management of obstetrics are used to manage pregnant women [8]. These guidelines stipulate that a pregnant woman should attend at least three antenatal visits during the first, second, and third trimesters. This allows monitoring of the health of both the fetus and the mother by healthcare professionals that includes midwives, obstetricians, and sonographers. During antenatal visits, pregnant women are referred for a medical ultrasound to monitor the well-being of the fetus and help with the early identification of fetal abnormalities, and where possible initiate relevant interventions.

To the best knowledge of the authors, this is the first case report on PUV to be documented from Botswana. At the time of this report, there has never been any known published cases of PUV from a medical imaging perspective. Most of the cases reported in the international literature on this topic are from medical professionals [1, 6, 7, 9]. We, therefore, report a rare diagnostic imaging case of PUV detected during a routine antenatal ultrasound in Botswana.
DIAGNOSTIC IMAGING CASE REPORT

A 22-year-old pregnant African ethnicity woman (gravida 3 para 2) presented in her fifth month to the outpatient department (OPD) for antenatal services in March 2021. By her dates, she was 20 weeks pregnant. As per procedure, she was referred to our imaging department by the attending obstetrician for her second-trimester prenatal ultrasound scan. An ALOKA 350 plus ultrasound machine with a 2.5-5.0 MHz transabdominal transducer was used to assess the pregnancy throughout this diagnostic imaging case report.

First ultrasound scan

The first prenatal routine ultrasound at 16 weeks was unremarkable and showed normal fetal kidneys, bladder, and adequate amniotic fluid volume (Figure 1). As per ultrasound guidelines, a second-trimester fetal anomaly scan was arranged at 20 weeks gestation age.

Second ultrasound scan

The second ultrasound revealed a male live fetus of 20 weeks gestation age, with bilateral hydronephrosis, distended urinary bladder, and significant oligohydramnios (Figure 2a). Figure 2b shows male fetal genitalia demonstrated during ultrasound scans. With the abnormal sonographic findings, a follow-up rescans at 24 weeks was arranged to assess the fetal urinary system.

Figure 1: 16 weeks gestation scan with normal fetal kidneys

Figure 2a: Bilateral fetal hydronephrosis at 20 weeks gestation age

Figure 2b: Fetal male genitalia at 20 weeks
On enquiry, her previous pregnancies were not associated with any fetal-maternal complications. The woman had no medical or surgical problem and there was no family history of chromosomal, congenital, or growth anomalies. She was generally in good health. 

**Third ultrasound scan**

A follow-up rescans at 24 weeks gestation age revealed a grossly distended urinary bladder, bilateral hydroureteronephrosis, and anhydramnios with reduced fetal movement (Figure 3).

![Figure 3: Bilateral fetal hydronephrosis at 24 weeks gestation age](image)

At this point, the sonographic findings raised a strong suspicion of PUV obstruction. The woman was counseled by the midwife and the attending obstetrician regarding the possible poor prognosis. The attending obstetrician discussed the fetal scan findings with the sonographer and a follow-up rescan at 27 weeks was arranged to monitor the fetal wellbeing.

**Fourth ultrasound scan**

The follow-up rescan was performed at 27 weeks and revealed a live fetus in breech presentation, reduced hydronephrosis compared to the previous scan, grossly over distended urinary bladder with generalized wall thickening (Figure 4), and anhydramnios.

![Figure 4: Over distended fetal urinary bladder at 27 weeks gestation age](image)

The neonatal specialist based at the referral hospital was informed and advised close monitoring with possible admission as a case of congenital PUV with poor prognosis. The woman was admitted and informed of the available clinical optional intervention of medical termination of the pregnancy, as newer modern interventions to improve the prognosis were not available in the country. Fetal growth continued in the 50th percentile with grossly reduced amniotic fluid volume. From this time, the pregnancy was managed as “high risk” and the woman was informed by the attending obstetrician of the possible medical termination or adverse pregnancy outcome.

**Fifth ultrasound scan**

An ultrasound scan follow-up a fortnight later was performed which continued to show anhydramnios, bilateral hydroureteronephrosis, over distended urinary bladder with a “keyhole sign” (Figure 6a), and reduced fetal movement. Both kidneys showed increased cortical echogenicity, with fullness of the central pelvis (Figure 6b).

Figure 6a: Urinary bladder showing a keyhole sign

Figure 6b: Increased renal cortical echogenicity

Given the persistent sonographic findings of anhydramios, overdistended urinary bladder, and bilateral hydroureteronephrosis with its associated fetal complications, the mother received continued support and counselling from the obstetrician. Due to multiple organ involvement, and continued poor prognosis, she was informed and advised to opt for medical termination of the pregnancy which she declined.

The outcome of the pregnancy

At 30 weeks of gestation age, spontaneous labour occurred, and a still fresh birth male fetus was delivered with a fetal weight of 1900g and a length of 43.1cm. The imaging department was informed of the outcome and immediately the whole-body plain X-ray often called ‘Baby-gram’ was requested by the obstetrician and performed to check for any missed possible skeletal abnormalities on prenatal ultrasound scans which appeared normal (Figure 7).

Figure 7: Whole-body X-ray (Baby-gram)
A thorough physical examination was done by the obstetrician on the fetus which showed a moderately distended urinary bladder. The male external genitalia were normally developed; a fine cannula could be passed through the fetal urethral meatus but could not be passed into the urinary bladder due to an obstruction at the level of the urethral vesical junction. The physical assessment and all sonographic findings proved this case as of PUV.

**DISCUSSION**

In this case report, medical ultrasound was used as the first-line imaging modality to diagnose PUV. Unlike plain film X-rays, ultrasound imaging is free of ionising radiation, and is regarded as the safest diagnostic tool for pregnant women [10, 11]. The World Health Organization (WHO) [12] recommends three antenatal ultrasound scans during pregnancy. The first scan at 8-12 weeks to confirm intrauterine pregnancy, the number of fetuses, gestation age, and early detection of abnormal pregnancy; the second scan at 16-20 weeks to assess for any fetal anomalies; the third scan at 32 weeks onwards to determine fetal presentation, placental location, estimation of fetal weight (EFW) and general fetal wellbeing. However, ultrasound is highly operator dependent [11]. This means that to have a correct diagnosis, the sonographer should be appropriately be trained and experienced. For this reason, the WHO [12] recommends that antenatal ultrasound be performed by a competent person to enable the detection of any pregnancy complications and facilitate timely and appropriate management. However, there is an acute scarcity of sonographers in Botswana. This can be attributed due to the non-availability of a training institution offering medical imaging courses, such as radiography and ultrasound.

Antenatal ultrasound plays an important role in the identification of fetal gender. This is of great importance in the management of pregnancy in families at increased risk of sex inherited diseases and diagnosis of fetal abnormalities such as PUV [11, 13]. This is best performed in the second trimester of pregnancy when the external genitalia are well developed and visualized, and the fetus can freely move in the amniotic fluid [13]. The gender diagnosis is made by visualization of the external genitalia: a boy is diagnosed if the sonographer sees the penis and scrotum and a girl if the labias are seen. Literature reports that PUV occurs exclusively in males due to the complex development of a male genital urinary system which occurs between the fourth and fifth week of embryogenic life [5-7]. This was as in our case where male fetal genitalia was confirmed by ultrasound at 20 weeks of gestation age (Figure 2b). In the context of the case under discussion, the sonographic finding of a male was the first feature that narrowed the diagnosis to PUV.

One of the complications of PUV is obstruction of the urinary tract affecting the kidneys, ureters, and urinary bladder [3, 6]. In this case report, there was bilateral hydroureretorenphrosis and a grossly thickened wall distended urinary bladder which resembled a keyhole. In ultrasound, this is referred to as a “Keyhole sign” [6, 7]. The obstruction of the fetal urethra resulted in the build-up of urine in the kidneys and ureters (hydroureretorenphrosis) and over-distension of the urinary bladder. This sonographic finding agrees with what is reported in the literature regarding PUV [3, 6, 7]. If no intervention is done, this may result in damage to the fetal kidneys, which may be ascertained sonographically as abnormal increased renal cortical echogenicity [7] (Figure 6b). This second sonographic feature of an obstructed urinary tract narrowed the diagnosis to that of the PUV case.

During pregnancy, the amniotic fluid provides necessary fluid, space, and growth factors to allow normal development and growth of fetal organs such as lungs [14, 15]. In this case, there was a significant reduction in amniotic fluid volume as the pregnancy progressed. Suliburska et al., [14] and that fetal urination is the main mechanism contributing to the amniotic fluid volume and its essential role in the normal development and maturation of the lungs. Its persistent deficiency may result in lung hypoplasia, which is the incomplete development of parts of the lungs [15]. Because of the gross oligohydramnios at 30 weeks, there was significantly reduced fetal movement. Ultrasound also revealed increased abnormal renal echogenicity which is consistent with fetal renal failure which has a high prenatal morbidity and mortality rate [16]. This third sonographic finding of oligohydramnios added to the evidence of a case of PUV.

Literature reports that PUV is associated with skeletal abnormalities such as craniospinal defects and poor bone growth [9]. To rule out skeletal abnormalities which could have been missed during an antenatal ultrasound, a ‘babygram ’was performed on the fetus. An anterior-posterior view of the whole body was imaged as recommended in the radiography literature [17]. The “babygram” revealed no evidence of associated skeletal abnormalities (Figure 7).

**CONCLUSION**

PUV is a rare congenital malformation affecting the urinary system in males. Medical ultrasound is regarded as the first-line imaging modality used to diagnose this fetal anomaly. This imaging case report demonstrates that prenatal diagnosis requires follow-up serial ultrasound scans which play an important role in the assessment and progression of the complications that come with the pregnancy. It is highly recommended to have a minimum of three prenatal ultrasound scans: in the first, second, and third trimesters. However, there is a scarcity of sonographers in Botswana to cover a comprehensive ultrasound service. To improve the quality of antenatal services, it is recommended to train or recruit more sonographers.
CONSENT FOR PUBLICATION

Consent was obtained from the patient to use the information from her scan results. Permission was also sought and granted by our hospital management to have access to the patient’s information and use sonographic and X-ray images in this publication [Ref: GDHMT; 4/7/61].

REFERENCES