Transient Posterior Urethral Valve: Sonographic Diagnosis (A Case Report)

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SUMMARY

The greatest yield of fetal anomalies discovered in utero by ultrasound has occurred in the genitourinary system. Several studies have documented striking increases in the bladder or upper collecting system, which have spontaneously resolved and have not been felt to represent true and permanent obstruction. In the present study a case of "transient posterior urethral valve" diagnosed by ultrasonography has been reported with a review of the literature.

Key Word*: Obstructive uropathy, posterior urethral valve, prenatal ultrasonographic diagnosis.

The ability to image fetal anatomic parts and the understanding of ultrasonographic fetal anatomy have improved as the result of developments in instrumentation. This information is of value in determining fetal prognosis, in deciding the method of delivery and in alerting the pediatric staff to supportive and corrective measures which may need to be taken in the postnatal period. Recent developments allow antenatal therapeutic interventions such as the relief of urethral obstruction by the intrauterine insertion of a self-retaining suprapubic catheter which provides a bladder-amniotic fluid shunt. Since this is an invasive procedure with yet unproved clinical efficacy (1, 2, 4, 5, 7, 8, 10, 11, 12).

The greatest yield of fetal anomalies discovered in utero by ultrasound has occurred in the genitourinary system. The most common condition encountered is obstructive uropathies. Since the side and extent of obstruction are so varied, it is difficult for caretakers to assess the prognosis and develop a plan of management for each patient (2, 5, 7, 8, 9, 10, 11). At present, stenosis of urethra is one of the congenital defects for which intrauterine intervention can be considered. However the degree and the time of onset of the obstruction vary widely, and the optimal time of intervention is currently not known (2, 3, 6, 7, 8, 9, 10).

In this paper we described a fetus in which an obstruction of the urethra was diagnosed at 30 weeks of gestation. Subsequently we were able to monitor the pregnancy sonographically until birth at 36 weeks without interference.

CASE REPORT

Because of clinical suspicion of intrauterine growth retardation a sonographic examination was performed in a 25 years old primigravid woman at 30 weeks of gestation. A distended fetal bladder, bilateral hydronephrosis and bilateral hydroureters were found (Figure 1, 2, 3). Since amniotic fluid volume was found normal, so severe urethral obstruction could not be assumed. Therefore we decided to monitor the pregnancy without interference. Until 34 weeks the subsequent weekly scans showed gross alterations from the first scan and the amniotic fluid
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Figure 1. Sagital section through fetal abdomen demonstrating hydronephrosis and distended bladder.

Figure 2. Ultrasonic appearance of left hydronephrotic kidney.

Figure 3. Oblique scan through fetal abdomen demonstrating hydronephrosis of right fetal kidney and hydrourerter.

Volume persisted to be normal. Eventually at 34 weeks of gestation we found out that hydronephrosis and hydroureters disappeared spontaneously. Labor ensued at 36 weeks and the patient was delivered of a 2250 gm. male infant without any deformities.

DISCUSSION

This case suggests that an obstruction due to an urethral valve may rupture spontaneously or the ball valve mechanism may be reserved by the change in dynamics associated with bladder drainage (3, 6, 7, 10, 11). This phenomenon may represent a fortuitous bonus to intrauterine shunting in some fetuses or may be an unrelated event. Moreover, spontaneous rupture could well occur in fetuses without surgical intervention (1, 3, 5, 6, 7, 10, 11, 12).

Since hydronephrosis be transient before contemplating interventional procedures in utero which could cause many complications, one should make sure that hydronephrosis is persistent and that there is a genuine reduction in amniotic fluid quantity. Obstruction can not be considered on all or nothing phenomenon and different grades of stenosis will give different appearances that vary from bladder dilatation alone to bladder dilatation with bilateral hydronephrosis.

One is unable to determine from ultrasound appearance whether irreversible renal changes have already occurred. Rigid rules can not be made on the basis of available published data. For the present, however, some basic judgements can be made concerning the prognosis for some of these variations and the risks and benefits must be carefully assessed before a plan of management is chosen.

REFERENCES