DETECTION OF URINARY TRACT ANOMALIES USING ULTRASONOGRAPHY DURING ANTENATAL SCREENING

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ABSTRACT

Background & Aim: To study the pattern and prevalence of urinary tract anomalies, which can be diagnosed by USG during the antenatal period and to look for any associated anomaly. Material & Method: This study was carried out on a total of 1750 patients in SSG Hospital Vadodara. All the patients had come for their routine ante-natal screening. All USGs were done using curvilinear low frequency (3 -5 MHz) transducers. Results: Out of the 1750 patients studied over a period of 8 months, 19 patients were found to be having urinary tract anomalies. The prevalence of urinary tract anomaly in our study was 10.86 per 1000 fetuses (1.08%). In our study of the 19 fetuses with urinary tract anomalies, 2 had unilateral absent kidney, 2 fetuses were of Cystic renal dysplasia, 2 had ectopic kidneys and 1 patient, each of PUJ obstruction, vesicoureteric reflux, megaureter and posterior urethral valve giving the typical key hole appearance of the bladder. There were 2 fetuses with bilateral absent kidneys, oligohydranmios and pulmonary hypoplasia (Potter's sequence) which had to be terminated. Associated anomalies detected were of prune belly syndrome in association with megaureter. There were 2 fetuses with bilateral echogenic kidneys associated with absent radius. A fetus with bilateral cystic renal dysplasia also had associated hemi-vertebra and pulmonary hypoplasia. In our study, 26 % of the patients reflected changes in liquor be it oligohydranmios or polyhydranmios and 21 % of the patients had to be terminated prematurely as the anomalies wouldn’t have sustained in the late gestation. Conclusion: Urinary tract anomalies can be precisely diagnosed and classified in the antenatal period using ultrasonography imaging and it can be helpful in planning immediate postnatal care.

Key-words: Congenital anomaly, Urinary tract anomaly, Hydronephrosis, Antenatal.

INTRODUCTION

Urinary tract anomaly is very common in occurrence, with the rate of 1.08 %, i.e. approximately 1 urinary tract anomaly is encountered in every 100 ante-natal patients scanned. A systematic approach to the prenatal diagnosis of urinary tract abnormalities includes assessment of amniotic fluid volume, localization and characterization of urinary tract abnormalities, assessment of fetal gender,
and search for associated abnormalities. Normal AFV in the second half of pregnancy implies at least one functioning kidney and a patent urinary conduit to the amniotic cavity. In the setting of a urinary tract abnormality, normal AFV indicate a good prognosis. Occasionally and paradoxically, polyhydramnios may occur, especially with unilateral obstructive uropathy, with meroblastic nephroma, or when there are concomitant abnormalities of the central nervous system (CNS) or gastrointestinal (GI) tract. A detailed anatomic scan to search for associated abnormalities, which may indicate the presence of a syndrome or chromosomal abnormality. Renal anomalies may be part of the VATER association (vertebral defects, anal atresia, trachea-esophageal fistula, radial defects and renal anomalies). An expansion of this syndrome, VACTERL, includes cardiac and non-radial limb defects. The present study was undertaken with the aim to study the pattern and prevalence of urinary tract anomalies, which can be diagnosed by USG during the antenatal period and to look for any associated anomaly.

MATERIAL & METHOD

This prospective longitudinal study was conducted in the Department of Radiology, SSG Hospital & Medical College, Baroda, Gujarat during the period of one year. All cases, which were brought to the department of Radiology for antenatal screening were taken into the consideration. Ethical clearance was obtained from the Institutional Ethical Committee before commencement of the study. All the patients who came for their routine ante-natal screening were examined by Ultrasonography using curvilinear low frequency (3-5 MHz) transducers after taking informed consent. Detailed examination was conducted and data were collected in a pre-designed proforma and analyzed after comparing with other authors.

RESULT

One thousand and fifty cases of pregnancy were screened out by USG abdomen to diagnose any urinary tract anomaly and out of them 19 patients were found to be having urinary tract anomalies. The prevalence of urinary tract anomaly in our study was 10.86 per 1000 births. The anomalies we encountered were ranging from simple ones like single kidney to complicated ones like mega-ureter in prune-belly syndrome and urinoma in a posterior urethral valve. The most common anomaly was dilation of the renal pelvis, which was compatible with life and didn’t require any intervention post-natally. Single absent, dysplastic and ectopic kidneys had uneventful course, while the bilaterally absent kidney was found to be lethal anomaly. VUR, PUJ obstruction, mega-ureter, urinoma, PUV and bilateral echogenic kidneys required intervention post-natally. 26 % of the patients reflected changes in liquor be it oligohydramnios or polyhydramnios. 21 % of the patients had to be terminated prematurely as the anomalies wouldn’t have sustained in the late gestation.

DISCUSSION

The prevalence of urinary tract anomaly in our study was 10.86 per 1000 fetuses (1.08%). In a study conducted in Denmark by Rasmussen et al, the prevalence of fetuses with renal anomalies at 2nd
trimester scans was found to be 11.4 per 1000 fetuses (1.14%). In our study of the 19 fetuses with urinary tract anomalies, 2 had unilateral absent kidney, 2 fetuses were of cystic renal dysplasia, 2 had ectopic kidneys and 1 patient each of PUJ obstruction. Vesicoureteric reflux, megaureter and posterior urethral valve giving the typical key hole appearance of the bladder. There were 2 fetuses with bilateral absent kidneys, oligohydramnios and pulmonary hypoplasia (Potter's sequence) which had to be terminated. Associated anomalies detected were of Prune Belly Syndrome in association with megaureter. There were 2 fetuses with bilateral echogenic kidneys associated with absent radius. A fetus with bilateral cystic renal dysplasia also had associated hemi-vertebra and pulmonary hypoplasia.

In our study, 26 % of the patients reflected changes in liquor be it oligohydramnios or polyhydramnios. 21 % of the patients had to be terminated prematurely as the anomalies wouldn’t have sustained in the late gestation, which is comparable to the data concluded by Rasmussen et al, where 127 of the 412 fetuses (30.8%) underwent termination and to the perinatal mortality as found by Taguchi K et al where 7 lethal anomalies were detected of the 56 fetuses (12.5%) with urinary tract anomalies.

1) UNILATERAL RENAL AGENESIS: Unilateral renal agenesis is three to four times more common than bilateral renal agenesis. It may be difficult to diagnose prenatally because AFV is normal and the bladder appears normal. If a kidney is not found in the renal fossa, most are either congenitally absent or ectopic. The contra-lateral kidney may be enlarged because of compensatory hypertrophy. (FIGURE 1).

2) BILATERAL RENAL AGENESIS: ultrasound findings include severe oligohydramnios and non-visualization of the kidneys and bladder. Distinctive, flattened appearance of the adrenal gland on longitudinal sonogram (“lying down” adrenal sign) helps to confirm that the kidney did not develop in the flank. Repeated and consistent, non-visualization of the urinary bladder (over 1 hour) is a secondary sign. Color Doppler ultrasound shows no renal artery
arising from the aorta (Au) in a fetus with bilateral renal agenesis.\textsuperscript{7} Before 16 weeks’ gestation, AFV is not dependent on urine production and may be normal despite the absent renal function. (FIGURE 2).

3) **RENAI\ L ECTOPIA**: Renal fossa is empty, careful scanning may demonstrate the ectopic kidney adjacent to the bladder or iliac wing. The ectopic kidney is located on the opposite side of the abdomen relative to its ureteral insertion into the bladder, resulting in crossed renal ectopia with or without fusion. In most cases the crossed kidney fuses with the normally located kidney (cross-fused ectopia), and an enlarged bilobed kidney is seen. Renal ectopia is associated with a high incidence of urologic abnormalities, most often VUR. (FIGURE 3).\textsuperscript{3}

4) **MCKD**: Malformed kidney is usually enlarged but may be normal or small. There are multiple cysts of varying sizes that do not communicate with each other and are randomly distributed. Large peripheral cysts distort the retiform contour. Pelvis and ureter are usually atretic and not visible. On color Doppler evaluation, the renal artery is either absent or very small.\textsuperscript{8} Occasionally, a MCDK with a large central cyst and small peripheral cysts can mimic hydro nephrosis from ureteropelvic junction (UPJ) obstruction. In hydro nephrosis, however, the dilated calyces are of uniform size and anatomically aligned and communicate with the dilated renal pelvis. The kidney usually maintains the reniform contour, with renal parenchyma present peripherally. On serial examinations, the kidney and its cysts may increase or decrease in size or may initially enlarge and later involute. (FIGURE 4).\textsuperscript{8}

5) **ARPKD**: Sonography reveals bilateral retiform enlargement of the kidneys. There is poor delineation of the intrarenal structures. The numerous tiny cysts are usually smaller than the limit of sonographic resolution, but they create multiple acoustic interfaces, accounting for the characteristic increased renal echogenicity and loss of corticomedullary differentiation. The kidney demonstrates increased echogenicity, with no visible cysts. There is loss of corticomedullary differentiation. When renal function is abnormal, there is oligohydramnios, and the bladder is small to absent. (FIGURE 5).\textsuperscript{12,13}

6) **BRIGHT KIDNEYS**: Sonographic evidence of urinary tract obstruction, renal dysplasia is a possibility, especially when the kidneys are small or normal in size and there are peripheral cortical cysts. Kidneys and the biometric measurements are above the 95th centile, an overgrowth syndrome (Beckwith-Wiedemann syndrome, Perlman syndrome) should be considered. In both Beckwith-Wiedemann syndrome, there may be macroglossia and omphalocele. In Perlman syndrome, there may be micrognathia and a depressed nasal bridge. Isolated, bilaterally enlarged hyperechogenic kidneys, the most common underlying diagnosis was ARPKD, followed by ADPKD (FIGURE 6).\textsuperscript{13}
7) **MesoBlastic Nephroma** is indistinguishable from Wilms’ tumor. Mesoblastic nephroma is usually seen as a moderately echogenic, solid mass completely replacing the kidney or localized to the part of the kidney. The mass may demonstrate increased vascularity and cystic components. Polyhydramnios is a frequent association and may lead to preterm labor and preterm birth. Perinatal complications are likely, including acute fetal distress, neonatal hypertension, and hypercalcemia. (FIGURE 7).

8) **PUJO:** Dilated renal pelvis with or without cablecasts is identified. The ureter and the bladder are not dilated. Severe chronic obstruction leads to the effacement of the calyces and thinning of the renal cortex (SFU grade 4). Rarely, the renal pelvis may be extremely dilated, presenting as a large, unilocular cystic mass. Rupture of the collecting system results in the development of a perirenal urinoma. (FIGURE 9). This “pop-off” mechanism may protect the obstructed kidney from further prenatal damage and may diminish the degree of hydronephrosis. The AFV is usually normal but may be increased paradoxically. When unilateral hydronephrosis is accom- panied by oligohydramnios, a search for contralateral renal pathology is warranted. (FIGURE 8).

9) **VUJO:** The affected kidney may demonstrate dilation of the ureter and renal pelvis. Serpiginous cystic segments must be traced to the renal pelvis and bladder. Megaureters are classified into three types per their morphologic appearance. Type I megaureter dilation involves only the distal ureter, with a normal-appearing upper tract. Type II extends to both ureters and pelvis. Type III is associated with severe hydronephrosis and ureteric tortuosity. To differentiate the dilated ureter from the bowel is important. Bowel contents are usually more echogenic than urine. In addition, the ureter generally comes into close contact with the spine, but the small bowel does not. (FIGURE 10).

10) **HN and VUR:** The most common prenatal sonographic diagnosis is hydronephrosis, which may be unilateral or bilateral. The ureter may be dilated. Intermittent dilation of the collecting system favors VUR. Fluctuation or variation in the RPD (changing by more than 3 mm) during an obstetric sonogram was strongly associated with high-grade VUR (grades IV-V). (FIGURE 11).

11) **Fetal megacystis** has been reported as early as 10 to 14 weeks gestation when the longitudinal bladder diameter is 7 mm or more. Severe megacystis (bladder length >15 mm) Posterior urethral valves are the most common cause of lower urinary tract obstruction, followed by urethral atresia or stricture. Posterior urethral valves are seen exclusively in males and may cause total, intermittent, or partial obstruction, with variable prognosis. Back pressure causes a persistently dilated urinary bladder, with a dilated proximal urethra (keyhole sign) (FIGURE 12).

12) **Prune Belly Syndrome** is characterized by the classic triad of
absent abdominal musculature, undescended testes, and urinary tract abnormalities (megacystis, ureterectasis). The bladder is typically very large. The prostatic urethra is dilated, and the appearance resembles posterior urethral valves. The ureters tend to be tortuous and dilated. The kidneys may be normal, hydronephrotic, or dysplastic. Other abnormalities may be present, including intestinal malrotation, congenital heart disease, and musculoskeletal deformities. (FIGURE 13).

Figure 1

Figure 2

Figure 3

Figure 4

Figure 5
CONCLUSIONS

The prevalence of different types of urinary tract anomalies is higher than that reported in developed countries. Urinary tract anomalies can be precisely diagnosed and classified in the antenatal period using ultrasonography imaging and it can be helpful in planning immediate postnatal care.

Source of Funding : Nil.
Conflict of Interest : None.

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