The Role of MRI in Evaluation of Fetal Urinary Tract Anomalies with Normal and Abnormal Volume Amniotic Fluid

Abstract

Purpose: to evaluate the role of magnetic resonance imaging (MRI) in evaluation of fetal urinary abnormalities with normal and abnormal volume amniotic fluid after ultrasound examination.

Methods: The prospective study from April 25, 2016 to January 21, 2017, comprised 16 fetuses with normal volume amniotic fluid 10 patients and abnormal volume amniotic fluid (an/oligohydramnios) 6 patients. All fetuses with suspected congenital urinary tract anomalies based on ultrasound underwent MRI examination with 1.5T and 3T MRI. The MRI findings were compared to ultrasound, micturition urethrocytography (MUC), computed tomography (CT) and magnetic resonance imaging (MRI) findings as a gold standard postpartum imaging and surgical findings. Gestational age range was 19-37 weeks.

Result: In anhydramnios 2 fetuses showed concordant with MRI (bilateral multicystic dysplastic kidney (MCDK) and normal kidney), 1 fetus with hypoplasia kidney between ultrasound (US) and MRI is discordance, in oligohydramnios 3 fetuses MRI has additional information (autosomal recessive polycystic kidney dysplasia (ARPKD) with fibrocystic liver, anterior urethral valve and posterior urethral valve obstruction). In normal amniotic fluid MRI gave additional information in 9 patients (left hydronephrosis (HN), prune belly syndrome, vesicoureteral junction obstruction (VUJO) bilateral, megaureter microcolon intestinal hypoperistaltic syndrome (MMIH), doaca, Left multicystic dysplastic kidney (MCDK), L pelvic cyst and horseshoe kidney), 1 patients showed concordant with MRI (L pelviuretero junction obstruction (PUJO)).

Conclusion: Fetal MRI can be used as a complementary tool in evaluation of fetal urinary tract anomalies with normal and abnormal volume amniotic fluid cases.

Keywords: Fetal MRI; Congenital urinary tract anomalies; Normal and abnormal volume amniotic fluid.

Introduction

Congenital urinary tract anomalies are relatively common, accounting +30-50% of all structural anomalies at birth [1]. The abnormalities are broad spectrum from mild to severe which is threatened of fetus. US is still using for screening urinary tract anomalies [2,3]. Some conditions such as anhydramnios and oligohydramnios, the results may be inconclusive. In these situations the organ cannot be evaluated sufficiently. It is a clearly indication to performed fetal MRI for urinary tract anomalies [4-8]. Otherwise in normal volume amniotic fluid urinary tract can be seen clearly by US. MRI has large field of view and good natural contrast can differentiate between dilated urinary tract and bowel and also determine level obstruction. The purpose of this paper is to evaluate the role of MRI in urinary tract anomalies with normal and abnormal volume amniotic fluid.

Materials and Methods

We prospectively examine 19 fetuses from April 25, 2016 to January 21, 2017 with suspected urinary tract anomalies based on ultrasound. Inclusion criteria presence of oligohydramnios or anhydramnios, and normal amniotic fluid with abnormal urinary tract or multisystem anomalies with improper visualization of kidney. MRI was done within 1 weeks following US. Three mothers of fetuses with contraindication to MRI and claustrophobia were excluded.

Sonography was performed by obstetrician using Samsung A30, full 2D combined gray scale and color Doppler studies and 3D, 3.5-5MHz. Amount of amniotic fluid using 4 quadrant amniotic fluid index. Color Doppler was done to search renal arteries in suspected renal agenesis.
MRI was performed with 1.5T Siemens Avanto, Erlanger, Germany, 1.5T GE, Signa HD, and Siemens Skyra 3T, using body coil with supine or LLD position. After a scout acquisition, a series of fetal images in axial, sagittal, and coronal T2 HASTE/FFSFE with 4 mm slice thickness and 30-35 cm FOV and sagittal T1 SPGR, heavily T2W MRU with 50 mm slice thickness, axial DWI b value 700. Contrast agent and sedation were not used. All MRI were analyzed by radiologists who experienced in fetal MRI who were blinded to US results. Amniotic fluid was qualitatively assessed as normal or diminished on the bases of the ability to identify at least three large pockets of fluid with a depth greater than 2 cm. Informed consent was obtained from all the mothers. Follow up was done for 10 fetuses with ultrasonography and contrast study (cystogram, retrograde pyelography, cloacogram), surgical exploration and CT, 6 fetuses were fetal demise and refused autopsy.

Results

Gestational age ranged from 19 to 35 weeks. Fetal movement did not alter image quality. We divided into 2 groups of patient between an/oligohydramnios (Table 1) and normal amniotic fluid (Table 2). In an/oligohydramnios diagnosis ultrasound revealed concordant with MRI 2/6 (33.3%) fetuses, 3/6 (50%) MRI gives additional information, 1/6 (16.6%) discordant (hypoplasia kidney). In discordant case USG revealed hydropsia but fetal MRI result agenesis. MRI was performed in 21 weeks gestation whereas intensity of kidney is similar with small bowel. Ultrasonography with color Doppler flow easier to visualize renal artery so more accurate to determine agenesis kidney. According to previous journal MRI is to be useful adjunct to US for the diagnosis agenesis kidney [5,9,10]. In this case MRI had no significant advantage over US unfortunately the parent refused autopsy (Figure 1). In case ARPKD US is sufficient to see enlargement of the kidney with hypoechoic in medullary kidney and fibrocystic liver (Figure 2), MRI has additional information for looking maturity of the lung. With increasing gestational age the intensity of the lung will increase as well [11,12]. In this case fetal MRI examination was performed in 35 weeks of gestation, but the lung intensity is still hypointens. An-oligohydramnios prevent growth thoracic cavity therefore maturity of the lung is inhibited which is cannot be seen by US. One case with bilateral hydroureretonephrosis and megacystic based on US, MRI gives additional information a cystic lesion on left kidney. Other case with hydroureretonephrosis and megacystis confirmed by MRI, MRI has additional information dilatation anterior urethra (Figure 3) was proven by MUC (Figure 4). One case with anhydramnios but normal both of kidney the result of US is the same with fetal MRI. The cause anhydramnios is premature rupture of the membrane.

Table 1: An/oligohydramnios.

<table>
<thead>
<tr>
<th>USG</th>
<th>MRI</th>
<th>Confirmation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilateral Renal Hypoplasia</td>
<td>Bilateral Renal Agenesis</td>
<td>Stillbirth</td>
</tr>
<tr>
<td>Bilateral MCDK</td>
<td>Bilateral MCDK</td>
<td>Stillbirth</td>
</tr>
<tr>
<td>Bilateral ARPKD, Fibrocystic Liver</td>
<td>Bilateral ARPKD, Fibrocystic Liver, Hypoplasia Lung</td>
<td>Stillbirth</td>
</tr>
<tr>
<td>HUN Bilateral, Megacystis</td>
<td>HUN Bilateral, Renal Cyst RK, Megacystis,</td>
<td>US, MUC</td>
</tr>
<tr>
<td>Normal Bilateral Kidney</td>
<td>Normal Bilateral Kidney</td>
<td>Stillbirth</td>
</tr>
<tr>
<td>HUN Bilateral, Megacystis</td>
<td>HUN Bilateral, Megacystis, Hydrourethra</td>
<td>MUC</td>
</tr>
</tbody>
</table>

Table 2: Normal amniotic fluid.

<table>
<thead>
<tr>
<th>USG</th>
<th>MRI</th>
<th>Method Confirmation</th>
<th>Final Diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>LHN</td>
<td>LHN</td>
<td>MUC, USG</td>
<td>VUJO</td>
</tr>
<tr>
<td>HN bilateral</td>
<td>Bilateral HUN Dominant on the Left Side</td>
<td>RPG at the Left Side</td>
<td>VUJO Left Side</td>
</tr>
<tr>
<td>Megacystis, Multiple Renal Cysts</td>
<td>Megacystis, Bilateral HUN, Cryptorchidism</td>
<td>MUC, Physical Examination</td>
<td>Prune Belly Syndrome</td>
</tr>
<tr>
<td>Megacystis, Keyhole</td>
<td>Megacystis, gastric dilatation, no meconium in large bowel, no keyhole</td>
<td>laparotomy</td>
<td>MMH</td>
</tr>
<tr>
<td>bilateral HN, Ascites, Multiple Pelvic Cysts</td>
<td>bilateral HUN, Hydrometrocolpos, Ascites, Abnormal Rectum Position</td>
<td>Cloacogram, Laparotomy</td>
<td>Cloaca Malformation</td>
</tr>
<tr>
<td>LHN</td>
<td>LHN</td>
<td>US</td>
<td>PUJO</td>
</tr>
<tr>
<td>Agenesis LK, Multiple Pelvic Cyst</td>
<td>MCDK LK, no Pelvic Cyst</td>
<td>MRI</td>
<td>MCDK</td>
</tr>
<tr>
<td>Megacystis, HUN Bilateral</td>
<td>HN RK, HUN LK with Ureneterocele and Cystic Dysplastic, No Bladder</td>
<td>N/A (stillbirth)</td>
<td></td>
</tr>
<tr>
<td>L Renal Cyst</td>
<td>L Pelvic Cyst with Calyceal Dilatation</td>
<td>USG</td>
<td>L Pelvic Cyst</td>
</tr>
<tr>
<td>Multiple Congenital Anomalies, Improper Visualization of Kidney</td>
<td>Horseshoe Kidney with L HN</td>
<td>USG, CT</td>
<td>Horseshoe Kidney, L HN</td>
</tr>
</tbody>
</table>
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Figure 1a: CDFU shows renal artery. 1b: MRI coronal T2HASTE. 1c: Axial T2HASTE reveals non visual kidney at renal fossa.

Figure 2a: US shows enlarged echogenic medullary with sparing cortex. 2b: shows fibrocystic of the liver. 2c: Coronal T2HASTE MRI shows hyperintens enlargement both of the kidney and fibrocystic liver. Lung is still hypointens inapropriate with age of gestation. MRI was performed in 35 weeks gestation.

Figure 3a: Coronal T2HASTE showing hydroureteronephrosis and thinning renal cortex. 3b: Axial T2 bilateral ureter are dilatation (arrow). 3c: Showing dilatation anterior urethral (arrow) and irregular posterior wall (double arrow).

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In normal amniotic fluid group (Table 2) MRI gives additional information in 9 patients, 1 patient with PUJO concordant between USG and MRI. Dilated collecting system is the most frequent cases 8/10 (80%) with variation causes: 2 cases due to VUJO, 1 case dilated collecting system due to prune belly syndrome, 1 case due to cloaca malformation, 1 case due to PUJO, 1 case horseshoe kidney, 1 case due to pelvic cyst, 1 case cannot be proved due to stillbirth. Others cases are megacystis microcolon hypoperistaltic syndrome (MMIH) without dilatation of collecting system and 1 cases MCDK of the left kidney.

In case of urinary tract obstruction US has limitation to see the whole length of the ureter for example US was diagnosed hydronephrosis, but MRI hydroureteronephrosis (Figure 5). In massif dilatation and tortuous ureter US has difficulties to differentiate between urinary tract and the bowel (Figure 6). In megacystis cases MRI can give definitive diagnosis as prune belly syndrome (Figure 7) and MMIH (Figure 8). In prune belly MRI reveals bilateral hydroureteronephrosis with cryptorchidism, megacystis with floppy shape bladder wall and dilatation posterior urethra. The floppy shape and absence of bladder wall thickening, cryptorchidism made the working diagnosis is prune belly rather than posterior urethral valve obstruction. MRI can give clear image of the bladder wall compare to US to differentiate between mechanical and functional obstruction. In mechanical obstruction bladder wall is thickened and irregular due to long standing obstruction. In MMIH MRI shows megacystis with dilated stomach, small large bowel and no meconium fills large bowel in female fetus. MMIH is a rare congenital syndrome. It represents severe form of functional obstruction due to abnormal action/myosin resulting hypoperistaltic of the bowel and urinary tract. Ultrasound cannot see the meconium so the diagnosis MMIH cannot be established by US.

In multiple pelvic cysts MRI can differentiate between bladder, genital tract and bowel. MRI can determine the level rectal termination which is cannot be seen by US (Figure 9). The depiction of multicycstic pelvic mass composed of bladder and hydrometrotcolpos, ascites, hydronephrosis and abnormal rectal position should lead to the diagnosis of cloaca malformation [13,14]. Ascites in this case is due to urine refluxing via fallopian tube. Hydronephrosis is occurred due to bladder obstruction by pouch. Cloacagram with contrast water soluble via single opening and colostomy confirm cloaca malformation (Figure 10).

In left MCDK case US was diagnosed agenesis left kidney due to small field of view (FOV) of the US and the cysts are large enough to occupy the lower abdomen making misdiagnosis as multiple pelvic cyst. Moreover in MCDK there is no normal renal parenchymal can be seen that could make difficulties to determine that the cysts are from the kidney (Figure 11). In megacystic and bilateral hydroureteronephrosis case MRI shows no bladder, right hydronephrosis, left hydroureteronephrosis, ascites and urinoma at the right kidney. Unfortunately the fetus is demised and refused autopsy, but karyotyping result trisomy 13 (Figure 12).
In multiple congenital anomalies with improper visualization of kidney MRI can give additional information the shape and dilated pelvicalyceal of the left kidney. These findings are in line with CT finding postpartum (Figure 13).

**Figure 6a:** MRI at 19 week gestation, coronal T2 HASTE MRI shows left hydroureteronephrosis. 6b: axial MRI shows the right pelvicalyceal dilatation as well (arrow). 6c: Follow up MRI at 32 weeks gestation showing right pelvicalyceal dilatation (arrow) and huge left ureter dilatation (asterix). 6d: RPG at the left side showing dilatation pelvicalyceal and ureter, no contrast fills bladder.

**Figure 7a:** Thick slab MRI showing hydroureteronephrosis bilateral and megacystis with dilated posterior urethra. 7b: axial T2HASTE reveals cryptorchidism making the first diagnosis is prune belly. 7c: MCU shows reflux contrast grade 4.

**Figure 8a:** Sagital T2HASTE showing megacystis. 8b: Axial T2HASTE shows small colon (arrow). 8c: enlargement of stomach (arrow). 8d: T1SPGR reveals no meconium in colon. 8e: Surgical exploration shows malposition microcolon (arrow head), meconium fill small bowel (arrow), no mechanical obstruction.
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Figure 9a: Coronal T2HASTE showing bilateral hydronephrosis and large pouch of vagina. 9b: Coronal T2HASTE reveals meconium level in urine containing pouch (arrow), bladder compressed by pouch (star). 9c: Sagittal T2HASTE Bladder compressed by pouch of hydrometrocolpos, sacral segment present (double arrow), no rectum between pouch and spine. 9d: T1SPGR shows high rectal termination.

Figure 10a: single opening. 10b: Cloacagram shows connection rectum with pouch. 10c: Cloacagram shows connection between vesica urinaria (VU) and pouch.

Figure 11a: MRI Coronal. 11b: axial T2HASTE reveals multiple variable sized non communicating cysts. 11c: Axial T2 HASTE post partum MRI shows L MCDK.
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Discussion

In clinical practice, the ultrasound is still the examination of choice for fetal imaging because of its availability in health facilities, affordable in financial side, and sufficient to answer most of clinical questions. But there are cases with subtle findings on US or clinical problems that US cannot nor hard to give explanation because of the nature of the examination and also because the maternal condition that give not enough windowing for the US. Fetal MRI is one of the advanced imaging that could help and acts as adjunct examination to answer those questions [14].

Our series found that the MRI on fetal urinary tract anomalies gave a more thorough and detailed anatomy without concerning about abnormally amniotic fluid volume (anhydramnios nor oligohydramnios). Even in cases with normal amniotic fluid, fetal MRI is able to give more information regarding the anomalies. With MRI, we can gain findings such as level obstruction in the case of hydronephrosis, the side severity in bilateral involvement of the urinary tract dilatation, association of other anomalies like cryptorchidism, and gastrointestinal abnormalities which contribute in a congenital non syndromic nor syndromic anomalies such as MMH (gastric dilatation, abnormal passage of the meconium), cloacal malformation, ureterocele, and also gives a clear images regarding anomalies of renal morphology (i.e. MCDK, horseshoe kidney) and more detailed anatomy on associated neural tube defect [15,16]. In cases with extraurinary involvement, MRI can detect intestinal anomalies by visualizing the exact part of the gastrointestinal that has the abnormalities by identifying the distribution of meconium within the bowel and its ability to differentiate between water and meconium [15]. In our cases with anhydramnios or oligohydramnios, most of the US findings in concordance with the MRI. The superiority of the fetal MR is the ability to see and predict the maturity of the lung parenchyma and lung volume that will affect the patient’s pre- and post-treatment [12]. The additional information provided by MRI will be one from many considerations in the decision of

Figure 12a: MRI T2 HASTE showing severe hydronephrosis with parenchymal thinning and urinoma (arrow). 12b: Sagital T2HASTE MRI shows ureterocele (arrow). 12c: axial T2HASTE reveals dilatation of left ureter (arrow) and displastic changes of parenchymal kidney.

Figure 13a: axial T2HASTE shows dilatation pyelocaliceal left kidney and isthmus between right and left kidney (arrow). 13b: Sagital T2MRI shows spinal defect. 13c: CT postpartum showing horseshoe kidney with dilatation of the left pyelocalyceal.
termination choice in cases with severe abnormality that has a high mortality rates. In renal agenesis cases, MR has limitation in differentiating the signal intensity between the kidney and bowel because such cases the renal fossae occupied by the bowel. For that circumstance, the US has advantages with its Doppler ability. Our finding is consistent with Noha et al reported [16].

The limitation of our studies is small number of fetuses that included in the study. Our fetal MRI cases relatively low since the examination is still new thing for the practicing clinicians, the access to fetal MR facilities is also confined to a very few hospitals, and also very expensive sophisticated examination. The stillbirth cases have no proof since the autopsy is not a compulsory to do in our country.

Conclusion

Our study shows the capability of MRI to give additional information in fetal urinary tract anomalies both in cases with normal and abnormally low amniotic fluid volume. Prenatal US has limitation due to small field of view, influenced by acoustic windowing, body habitus of the patient and operator dependent. Fetal MRI has the advantage to give detailed anatomy and associated anomalies that change the management of the patient, pre and postnatal.

Acknowledgement

None.

Conflict of Interest

None.

References