Review

Congenital obstructive uropathy – Diagnostics for optimal treatment

Christian Radmayr

Department of Paediatric Urology, Medical University Innsbruck, Austria

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Abstract

Antenatal ultrasound screening techniques reveal urological abnormalities in about 1 of 500 cases with half of which belonging to hydronephrosis. Postnatal appropriate diagnostic modalities are obligatory for proper diagnosis. Nowadays, ultrasound is definitively one of the most useful imaging modalities for the upper tract in children with hydronephrosis. It is non-invasive, radiation free and can be repeated. In addition, ultrasound also offers excellent images of the lower urinary tract in the diagnosis of posterior urethral valves or ureteroceles for example.

An additional tool is isotope renal scintigraphy which is method to analyse differential renal function and drainage of the kidney as well as functioning cortical mass with an accurate image of renal parenchyma. Today MRI studies are becoming more and more popular. This technique is particularly suited to urological imaging, because of its excellent delineation of water/urine-containing structures.

The micturating cystourethrogram is still the gold standard in imaging the bladder and the urethra. It also serves as a functional investigation while the patient is voiding. The technique consists of catheterizing the person in order to fill the bladder with a radiocontrast agent and is, therefore, an invasive method.

Since there is a risk of an increase in severity and functional deterioration with time on the one hand but also improvement and stable function on the other hand, suitable diagnosis is mandatory in order to rule out the children at risk.

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Natural history of prenatally diagnosed hydronephrosis

Routine antenatal ultrasound screening was introduced 30 years ago, and rapidly gained wide acceptance in some countries. Urological abnormalities are detected in 1 in 500 pregnancies [1,2], among which half are unilateral hydronephrosis (Fig. 1).

Before the era of prenatal ultrasound, hydronephrosis was mainly diagnosed in symptomatic older children presenting with loin pain, stones, haematuria, or pyelonephritis. Antenatal diagnosis has brought into light a unique and unknown population of healthy and asymptomatic infants having mild to severe hydronephrosis, with variable outcome [3].
It is known that prenatally diagnosed hydronephrosis may increase in severity and show functional deterioration with time, consistent with the definition of obstruction. It has also been reported that such a dilatation may remain stable (in fact the majority of cases), or improve with time, although it is unknown whether a true obstruction was present and vanished, or if the dilated but non obstructed system simply improved [4,5]. The potential for spontaneous resolution of unilateral hydronephrosis thus questions the necessity of surgical treatment, and the true nature of this dilatation if not obstruction.

Tribute must be paid to HK Dhillon and the perinatal urologists team of the Great Ormond Street Hospital for the outstanding amount of work dedicated to the follow-up of prenatally diagnosed hydronephrosis over a 25 years period. In their “natural history” series with conservative management, they showed that:

- Follow-up based on ultrasound is safe.
- Virtually all infants requiring pyeloplasty [6,7] for impaired renal function at birth had pelvic AP diameter >20 mm (Fig. 2).
- The vast majority of children born with normally functioning kidney, but who required surgery for deteriorating function had pelvic AP diameter >20 mm at birth.
- When deterioration of the function occurs, it is usually preceded by a worsening of the dilatation.

The same authors showed that ultrasound is the most useful imaging study to differentiate infants with PUJ obstruction who (will) require surgery from those who have pelvi-calyceal dilatation of no clinical significance. Among infants with unilateral hydronephrosis and normally functioning kidney on the postnatal MAG-3 renal scan, 90% of those with renal pelvis between 30 and 40 mm, and virtually all of those with renal pelvis >40 mm will eventually deteriorate. In contrast, among those with hydronephrosis <20 mm at birth, only 11% will eventually require surgery (they usually have major calyceal involvement, intra-renial pelvis, and often early perinatal diagnosis during the second trimester). The “grey zone” lies between 20 and 30 mm, where 40% may eventually deteriorate, but the same proportion may also improve spontaneously.

Other authors have also studied the natural history of prenatally diagnosed hydronephrosis with regard to the potential deterioration of renal function [8–11]. Koff followed non-operatively 104 neonates with unilateral hydronephrosis regardless of the degree of dilatation and of the initial degree of functional impairment. A large proportion of those with initial impairment of DRF rapidly improved spontaneously. Twenty-three of 104 (22%) ultimately required surgery for deterioration of either DRF, dilatation, or both, always before 18 months of age. Of those requiring pyeloplasty, there was no permanent loss of renal function and DRF eventually returned to exceed predeterioration levels.

Natural history series have provided very useful informations regarding to prenatal counselling and postnatal management strategy. The main points arising from these natural history studies are

- The majority of prenatual unilateral hydronephrosis (75%) will either remain stable or improve spontaneously
- Some of them will show increase of dilatation and/or deterioration of the renal function, and will ultimately require surgery [12–14]
- It is probably safe to closely follow-up dilated kidneys with ultrasound alone, and perform isotope imaging if hydronephrosis worsens
- The risk of deterioration correlates with the degree of dilatation of both pelvis and calyces. The threshold above which the risk of deterioration during follow-up seems unreasonable is about 30 mm.
- When normally functioning kidneys deteriorate during follow-up, many of them will recover preoperative function level after pyeloplasty.
- Management of kidneys with initial functional impairment is debated. Some of them may see their DRF improve spontaneously with time. In contradiction, even when surgery is performed, this subgroup of congenitally damaged kidneys seem not to improve as much after pyeloplasty as the ones with sudden deterioration. It is probably safe (and easier to follow-up) to perform pyeloplasty in infants with initial renal differential function impairment, although the renal outcome may not be different.

**Symptomatic hydronephrosis**

Symptomatic hydronephrosis has long been the main clinical presentation leading to the diagnosis of PUJ obstruction. Different complication could be observed, like febrile UTI, symptomatic stones, or intermittent abdominal pain. In a number of cases, the latter takes part of a very specific clinical picture named “acute intermittent hydronephrosis” [15,16]. Typically, these children (usually
old children or adolescents) will present with intermittent abdominal and loin pain, vomiting, and intermittent hydronephrosis usually related to lower-pole crossing vessels. It is important to note that ultrasound may display considerable variability of dilatation with time, and may be strictly normal between the crises (Fig. 3a and b). The frequency of the crisis will increase progressively if the diagnosis is not recognised. The majority of these children are thought to have no intrinsic PUJ obstruction.

Preoperative workup including US (during and out of crisis) studies MAG-3 functional assessment, but an angio-MRI may also be useful to assess the presence of crossing vessels, if not seen on ultrasound.

### Imaging studies

#### Ultrasound

Renal ultrasound is the cornerstone in the diagnostic pathway of children with hydronephrosis. During the prenatal period, one useful and reproducible method to assess hydronephrosis uses the Society for Foetal Urology (SFU) criteria [17], which grade the renal pelvic dilatation, the number of calyces seen and parenchymal atrophy within five grades of increasing severity (Fig. 4).

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No hydro</td>
</tr>
<tr>
<td>I</td>
<td>Renal pelvis is visualized</td>
</tr>
<tr>
<td>II</td>
<td>Renal pelvis and some calyces are visualized</td>
</tr>
<tr>
<td>III</td>
<td>Renal pelvis and all calyces are visualized</td>
</tr>
<tr>
<td>IV</td>
<td>Grade III plus parenchymal thinning</td>
</tr>
</tbody>
</table>

**Figure 4** SFU Grading System.

The degree of dilatation of the calyces, their size and shape, although more subjective, also appear very important in the analysis of the degree of dilatation. Knowing the anatomical variant of extra-renal dilated (but non obstructed) pelvis with normal calyces and parenchyma, we tend to define hydronephrosis as a dilated pelvis with some calyceal dilatation. Other criteria, like calyces/parenchymal thickness ratio, have proven less effective in determining the degree of dilatation.

In addition, ultrasound also offers excellent images of the lower urinary tract in the diagnosis of posterior urethral valves (Fig. 5) or ureteroceles (Fig. 6) for example.

#### Isotope renography: MAG-3 Renal scan

Dynamic diuretic isotope renal scintigraphy is a non-invasive method to analyse differential renal function and drainage of the kidney [18]. The radiopharmaceutical agent of choice is the 99mTc-mercaptoacetyltriglycerine (MAG-3), especially in infants: it is a tubular agent, with high protein binging, high tubular extraction, and low distribution in the extra-vascular space and subsequently low background activity. Serial dynamic acquisition show two distinct phases on a time-activity curve: an uptake phase with a rapid rise of activity as the tracer is delivered to the kidney, and a clearance phase as the tracer is excreted in the collecting system (Fig. 7).

#### Differential renal function (DRF)

Isotope renography provides a quantification of the differential renal function by comparing the slope of the time-activity curves of each kidney during the uptake phase. MAG-3 renal scan may also provide...
an estimate of the absolute renal function, although much less reliable than a glomerular filtration rate study using $^{51}$Cr-EDTA. In the assessment of unilateral uropathies, one has to keep in mind that DRF is a comparative analysis, even though the contralateral kidney is not normal. Hence, equilibrate split function does not always mean normally functioning kidneys, but sometimes equally impaired kidneys [19]. In addition, it is admitted that “static” DMSA renal scan provides a more accurate evaluation of the differential renal function especially with poorly functioning kidneys, whilst the irradiation dose delivered is also significantly higher.

**Clearance phase**

The decrease of activity in the region of interest, also called “drainage curve”, traduces the urine washout though the collecting system. There has been considerable debate regarding whether poor washout truly indicated the presence of obstruction in a dilated collecting system, or only delayed emptying. There are indeed several parameters involved, among which the degree of dilatation, the degree of maturation of the kidney, the degree of hydration, individual kidney function, urine flow rate, the timing of administration of the diuretic, the fullness of the bladder, and the timing of post-micturition images. It is now widely admitted that only looking at the slope of the drainage curve is a simplistic approach to a complex parameter.
The most obvious criterion is the dilatation: one cannot expect a large dilated pelvis to drain as quickly as a smaller volume pelvis [20]. It has also been shown that a full bladder will inhibit drainage from the pelvis, as well as the supine position; no conclusion on the drainage should be drawn without a post-micturition image and a change in posture.

Objective measurements, like half-time after diuretic injection, output efficiency, normalised renal activity, or pelvic excretion efficiency before and after voiding, have not proven more effective to assert PUJ obstruction. Studies looking at the natural history of unilateral hydronephrosis have reported several examples of impaired drainage curves in children whose dilatation and differential function remained stable over time, with no need for intervention (i.e. meeting the definition criteria of no obstruction).

In conclusion, there is little doubt that impaired drainage on diuretic renography is not synonymous of an obstruction in unilateral hydronephrosis, but may only reflect delayed emptying of a dilated system [8]. In contrast, good drainage on renography is surely a definite sign of the absence of obstruction.
Isotope renography: DMSA scan

Static renal scintigraphy is a non-invasive delivering images representing functioning cortical mass and therefore providing an accurate image of renal parenchyma [9]. The tracer bind to the proximal tubules and offers a high sensitivity in the detection of parenchymal pathology (Fig. 8).

Magnetic resonance imaging (MRI)

MRI is particularly suited to urological imaging, because of its excellent delineation of water/urine-containing structures. There has been great expectation that dynamic contrast enhanced MRI would be able to substitute to both renal ultrasonography and scintigraphy. Indeed, MRI has the ability to provide very detailed anatomical description (Fig. 9), combined with a functional evaluation. This promising technique, which has been under evaluation for more than 10 years, still struggles to provide functional informations as accurate as conventional isotope studies. Additional drawbacks are the costs, and the need for sedation or general anaesthesia after 6 months of age.

Pressure-flow studies

Whitaker first described this testing in the early 70s, which involved percutaneous nephrostomy, infusion of extrinsic flow (10 mL/min) and monitoring of intrapelvic pressures [21]. Not only this test was invasive, required general anaesthesia, and simultaneous monitoring with ultrasonography and fluoroscopy, but also appeared poorly discriminant and reproducible.
Nevertheless, refinements in the techniques of pressure flow studies allowed to show that renal pelvic pressures could remain normal at low infusion flow even in severely obstructed systems, giving support to the intuitive clinical impression that some collecting systems can remain equilibrated although obstructed, and that increased workload may lead to decompensation of pelvic pressures to dangerous levels [22].

Antegrade pyelogram

Percutaneous US guided puncture with antegrade pyelogram is rarely necessary but may be helpful in some confusing cases, when there is a doubt about the exact level of obstruction (Fig. 10).

Voiding Cystourethrogram

This radiologic methodology (also micturating cystourethrogram – MCUG), is still the gold standard in imaging the bladder and the urethra (Fig. 11). It also serves as a functional investigation while the patient is voiding (Fig. 12). The technique consists of catheterizing the person in order to fill the bladder with a radiocontrast agent [23–25] (Fig. 13). Under fluoroscopy (real time X-rays) the investigator watches the contrast enter the bladder and looks at the anatomy. If the contrast moves into the ureters and back into the kidneys, the diagnosis of vesicoureteral reflex is made. The degree of severity can be given ranging from grade I to V according to the scoring system of the International Reflux Study Group (Fig. 14). Additionally, it gives excellent information on duplex systems as well (Fig. 15a and b). The exam ends when the patient voids while the while watching under fluoroscopy allowing to get exact images of the urethra. It is important to watch the contrast during voiding, because this is when the bladder has the most pressure, and it is most likely this is when reflux will occur.

**Proposed pathway for prenatally diagnosed unilateral hydronephrosis**

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**Impaired differential function**

- **DRF <40%**
  - _pyeloplasty_
  - US at 3 months postop.
  - US + MAG-3 at 12 months postop.

- **DRF <10-15%**
  - repeat MAG-3 at 9-12 months
  - _DRF improved_
  - _follow-up_
  - nephrectomy systematic only if complication ?
Normal differential function

- **pelvis AP diameter 15-20mm**
  - if major calyceal dilatation, or early prenatal diagnosis
  - follow-up
  - US at 1 year
  - US at 2 years
  - US at 5 years
  - then discharge

- **pelvis AP diameter 20-30mm**
  - follow-up
  - US at 6 months
  - US + MAG-3 at 2 year
  - US at 2 years
  - US at 3 years
  - US at 4 years
  - US at 5 years
  - US at 7 years
  - US at 10 years
  - US at 15 years
  - ... then discharge

- **pelvis AP diameter > 30mm**
  - pyeloplasty

If increase hydroperosis: **MAG-3 renal scan**
If deterioration of DRF: proceed to pyeloplasty

DRF: differential renal function on isotope renography.
US: ultrasound renal scan.
AP diameter: antero-posterior diameter

**Conflict of interest**
None declared.

**References**


