PRENATAL INTERVENTION FOR URINARY OBSTRUCTION AND MYELOMENINGOCELE

HUBERT S. SWANA, RONALD S. SUTHERLAND, LAURENCE BASKIN

Department of Urology, University of California San Francisco, and Department of Surgery, Tripler Army Medical Center, San Francisco, California, USA

ABSTRACT

Widespread use of ultrasonography has resulted in an increase in the recognition of fetal hydronephrosis. The enthusiasm that accompanied early interventions has been tempered by the experience and results obtained over the past 2 decades. The goal has remained the same: to identify patients with serious prenatal obstruction and to identify those which may benefit from intervention. Myelomeningocele remains a devastating congenital anomaly. Fetal and experimental studies suggested that patients with myelomeningocele could benefit from prenatal intervention. Advances in technology and perinatal management have made intervention for more complex malformations such as myelomeningocele possible. This article will review current knowledge and will detail rational management for the management of prenatal hydronephrosis. The current state of antenatal myelomeningocele repair and the urologic implications will be described as well.

Key words: fetus; congenital abnormalities; prenatal diagnosis; myelomeningocele; intrauterine; surgery; fetoscopy

Int Braz J Urol. 2004; 30: 40-8

INTRODUCTION

During the past 2 decades pediatric urologists have begun acquiring patients with antenatally detected conditions. With the widespread use of maternal ultrasound, fetal hydronephrosis has become increasingly detected, and it comprises the most common prenatally diagnosed malformation. The concept of the unborn child as a potential surgical patient has become firmly established (1). Fetal medicine has rapidly evolved since early experiences with the management of fetal hydronephrosis. With time, the natural history and pathophysiology of urinary tract obstruction has become better understood. Improvements in diagnostic imaging tools, advances in fetal urine sampling, enhanced interventional techniques and equipment, and a better understanding of the risks and outcomes in these babies have helped to develop rational intervention and observation strategies. Nevertheless, the management of the fetus with hydronephrosis has remained controversial.

The purpose of this article is to review antenatal intervention and its history. The basics of normal fetal development will be integrated with the techniques used to diagnose disorders of the urinary tract. Particular attention will be devoted to the diagnostic techniques of ultrasound, fetal urine sampling and amniocentesis. Newer modalities such as fetal magnetic resonance imaging (MRI) will be described. Intervention for patients with myelomeningocele and the implications for urinary tract function will be discussed. Methods of intervention will be described along with their indications, contraindications, and complications.

SPECTRUM OF ANTENATAL DISORDERS: EMBRYOLOGY AND PATHOPHYSIOLOGY

Perturbation of the developing ureteral bud and its intended target, the metanephric blastema by
Prenatal Intervention

Distal obstruction affects normal renal development (2-7). By the 5th week of gestation, the ureteral bud rises from the mesonephric duct. It then begins to lengthen and canalize. Induction of the metanephric blastema occurs by the end of week 7. Primitive renal function begins between week 7 and 9, and by week 20, about 1/3 of the total number of nephrons are present. Nephrogenesis is complete by the 32nd week of fetal life, after which no demonstrable increase in the number of glomeruli is noted (8-10).

The spectrum of deleterious changes seen in antenatal urinary obstruction is the result of multiple factors. They include the time of onset, duration, and degree of urinary obstruction. In general the earlier the obstruction occurs the more disturbed the development of the fetal kidney (8). Renal dysplasia, the most severe form of renal injury and maldevelopment, has been attributed to a very early effect of elevated pressures in the urinary (2) and alternatively by ureteral bud malposition with subsequent misconnection between the bud and the metanephric blastema (4,11). Without ureteral bud induction, the blastema fails to develop. One sees clusters of disorganized metanephric structures surrounded by abundant fibrous tissue. Ninety per cent of cases of renal dysplasia are associated with urinary obstruction during nephrogenesis. Sonography is highly specific for diagnosing dysplasia and the demonstration of renal cysts in a fetus with known obstructive uropathy effectively indicates the presence of dysplasia (12). The absence of cortical cysts, however, does not exclude renal dysplasia.

Dilation of the urinary tract can be due to, ureteropelvic junction obstruction (UPJO), congenital obstructed and nonobstructed megaureter, multicystic kidney, duplication anomalies with upper pole ectopia or obstructing ureterocele, and vesicoureteral reflux (VUR) (13). Obstruction of the upper urinary tract from physiologic ureterectasis or from UPJO is rarely complete. One must also exclude physiological hydrenephrosis which usually spontaneously resolves prior to delivery or within the first year of life (14) (Table-1).

More distal causes of obstruction include posterior urethral valves, urethral atresia, cloacal anomalies, and prolapsing, obstructing ureteroceles. These entities can result in marked distortion of both ureters and kidneys as well as pathological bladder changes. The prune belly syndrome is rarely been associated with renal obstruction even though the urinary tract is massively dilated. Some have argued that the characteristic urinary tract dilation is a consequence of transient fetal urethral obstruction (15).

Spinal cord and subsequent vertebral formation begins at day 18 of gestation. Neural tube infolding (neurulation) occurs between 18 and 27 days of gestation and is normally followed by migration of mesodermal tissue around the developing spinal cord. The mesoderm gives rise to the vertebral arches, as well as the spinal and back musculature. The location, timing and extent of the abnormal closure lead to the varying degrees and levels of neural tube defects. Lesions can vary to include spina bifida occulta, (a closed tube defect), meningocele, (a protruding meningeal sac without neural elements), myelomeningocele (a meningeal sac with neural elements) and lipomeningocele (a meningeal sac with neural elements and fatty tissue). Myelomeningocele is the most common neural tube defect. Lumbar vertebrae are most commonly involved followed by sacral, thoracic and cervical vertebrae in decreasing frequency. Failure to close at the caudal end results in a distal defect with resultant lower limb paralysis and bladder dysfunction (16). An Arnold-Chiari type II malformation occurs in up to 85% of children with myelomeningocele (MMC). There can be herniation of the cerebellar tonsils through the foramen magnum. This can result in obstruction of the fourth ventricle and necessitates ventriculoperitoneal shunting.

Table 1 – Causes of prenatal hydronephrosis

<table>
<thead>
<tr>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ureteropelvic Junction Obstruction</td>
</tr>
<tr>
<td>Multicystic Dysplastic Kidney</td>
</tr>
<tr>
<td>Megaureter</td>
</tr>
<tr>
<td>Vesicoureteral Reflux</td>
</tr>
<tr>
<td>Ureteral Ectopia</td>
</tr>
<tr>
<td>Ureterocele Ectopic</td>
</tr>
<tr>
<td>Prune Belly Syndrome</td>
</tr>
<tr>
<td>Posterior Urethral Valves</td>
</tr>
<tr>
<td>Urethral Atresia</td>
</tr>
<tr>
<td>Pelvic Tumor</td>
</tr>
<tr>
<td>Cloacal Anomaly</td>
</tr>
</tbody>
</table>
Urology morbidity in patients with MMC is significant. Myelodysplasia can result in a poorly compliant bladder, sphincteric dysfunction, secondary vesicoureteral reflux, a predisposition to urinary tract infections, possible renal scarring and renal failure (17). Urologic morbidity is the sequela of neurologic injury. The neurologic deficit seen in MMC is believed to be due to several factors. The first is defective development. Evidence supporting a secondary insult to the exposed spinal cord has resulted in a “two hit hypothesis”. Histologic findings support the idea that the exposed spinal cord is vulnerable to damage by physical trauma as the cord contacts the uterine wall. Physical trauma, and the toxic effects of amniotic fluid and meconium to the exposed spinal cord have been reported (18-20). Fetal lower limb movements have been described in fetuses with MMC at 16-17 weeks (21). Animal studies, in which laminectomy was performed at mid-gestation, compared in-utero repair to no treatment. The animals that underwent fetal intervention were spared flaccid paralysis and incontinence of urine and stool (22). Histologic specimens of bladder tissue from children with spina bifida reveal increased intracellular matrix between muscle bundles, decrease muscarinic receptor density abnormal smooth muscle growth, and decreased innervation (23-25). These factors likely contribute to bladder dysfunction in human spina bifida patients.

**DIAGNOSIS OF OBSTRUCTION**

**Ultrasound**

The evolution of fetal intervention has paralleled the advancements in ultrasound technology. High resolution, real time imaging and the ability to choose focal zone depth have been major advances in ultrasonography (26). Fetal positioning plays a critical role in the interpretation and understanding of the fetal anatomy. The prone fetus is in the optimal position for imaging the kidneys (1). While the kidneys can be seen as early as the 15th week reliable imaging is not possible until week 18 (26).

Hydronephrosis is the most common cause of an abdominal mass in the neonate, and antenatal sonography readily detects fetal urinary tract dilation (1). Pelviectasis is found in 18% of normal fetuses (27). Both caliectasis and an anteroposterior pelvic diameter of greater than 10 mm have been proven to be reliable predictor of fetuses in need of postnatal urologic evaluation (28,29). The Society of Fetal Urology has adopted a grading system for hydronephrosis, which is widely used by pediatric urologists today (Table-2).

Prenatal sonography is very sensitive in differentiating ureteropelvic junction obstruction from other causes of obstruction and dilation (30,31).

While ultrasonography remains the primary imaging modality for the screening and evaluation of congenital abnormalities, it is not without limitations. Maternal obesity, oligohydramnios and suboptimal fetal position can make accurate imaging difficult. Early use of MRI was limited by slow acquisition times and was hampered by fetal motion. Newer methods have been developed that can reduce acquisition times and provide excellent image quality without the need for fetal sedation or paralysis (32). MRI can provide images unaffected by fetal position, maternal obesity, oligohydramnics, or overlying bowel and

---

**Table 2 – Ultrasound grading scale of hydronephrosis**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Central Renal Complex</th>
<th>Renal Parenchymal Thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Intact</td>
<td>Normal</td>
</tr>
<tr>
<td>1</td>
<td>Slight splitting</td>
<td>Normal</td>
</tr>
<tr>
<td>2</td>
<td>Evident splitting confined within renal border</td>
<td>Normal</td>
</tr>
<tr>
<td>3</td>
<td>Wide splitting pelvis outside renal border. Calices uniformly dilated</td>
<td>Normal</td>
</tr>
<tr>
<td>4</td>
<td>Further dilation of renal pelvis and calices</td>
<td>Thin</td>
</tr>
</tbody>
</table>

possibly could provide a definitive diagnosis of obstructive uropathy (33,34).

MRI seems to be superior in identifying the intracranial lesions such as agenesis of the corpus callosum, cerebellar dysplasia and holoprosencephaly that can accompany myelomeningocele (35). In the future, clinical decisions may be based on analysis of chemical and molecular events with MRI (36). Presently, MRI is a useful adjunct to ultrasonography. MRI provides additional information in myelomeningocele, other complex fetal cases, and cases of hydrenephrosis with indeterminate US studies.

**Fetal Urine and Amniotic Fluid Testing**

Invasive acquisition of fetal urine for analysis has become one of the most important measures of assessing fetal renal function. Measurement of fetal urine electrolytes and urinary proteins is a useful guide to the clinician in deciding whether prenatal intervention is indicated. Additional methods of assessing the overall status of the fetus include amniocentesis, chorionic villus sampling, percutaneous umbilical blood sampling (all for karyotyping), as well as amniotic fluid volume and its biochemical constituent measurement.

Fetal urine is normally hypotonic reflecting developing glomerular and tubular function (37). The amniotic fluid, in comparison, is somewhat hypertonic, and is not as reliable an index of renal function as the fetal urine. Determination of human fetal renal function is limited to simple concentration of specific urinary constituents. More physiologic measurements of glomerular function, while possible, are not routinely performed (38). Clearance of iothalamate has been done and shown to be non-predictive of renal outcome (10). Retrospective analysis of individual urine constituents have shown that a sodium of less than 100 mEq/L, osmolality less than 210 mOsm/L and chloride less than 90 mEq/L, if accompanied by lack of ultrasonographic evidence of dysplasia, are helpful in predicting residual fetal renal function. By categorizing patients according prognosis, assessment of the potential efficacy of intervention can be made (39). Elder et al. (39) and Johnson et al. (40) separately suggested that single determinations of urinary electrolytes may not be useful. Johnson et al. (41) proposed providing transient relief of obstruction by vesicocentesis followed by sequential sampling (3 or 4 samples over several days) of urinary electrolytes. This was felt to provide an assessment of the severity of the renal injury and potential for reversibility of renal injury. Those fetuses that experience an improvement in their biochemical parameters following decompression may benefit most from interventional therapy (41). Others have suggested that sampling of fetal urine electrolytes and osmolarity is not an optimal method to evaluate fetal renal function and recommend continued search for a better substance (42-44). A serum marker, which has provided some clinical utility, is beta-2 microglobulin. It is excreted by the kidney without placental cross-over so that fetal levels represent fetal renal function. One can see an elevation in renal dysplasia (45). Other urinary constituents commonly associated with the presence of renal disease include proteins such as albumin, retinol binding protein, and N-acetyl-b-glucosaminidase have been studied (45). Unfortunately human fetal urine sampling lacks known control normals at different stages of development. Further limitation includes the inability to accurately and physiologically measure renal function by fractional excretion of biochemical constituents and glomerular filtration without risky invasive fetal and maternal testing (46).

**INTERVENTIONAL TECHNIQUE**

**Current Indications and Contraindications**

For most fetuses with obstructive uropathy, intervention is not necessary (47) (Figure-1). The selection criteria for fetal therapy of obstruction evolved such that patient selection is presently good enough to avoid intervention in patients who are either too well (no benefit) or too ill to recover (48). It has been conclusively shown that decompression in utero will restore amniotic fluid, which can prevent the development of fatal pulmonary hypoplasia. What seems less clear is whether or not in utero decompression can arrest or reverse cystic dysplastic changes caused by obstruction (48).

Spontaneous resolution of hydrenephrosis is common, which has led to a more cautious approach.
to fetal intervention (1,49). In most cases with normal amniotic fluid volume, the mother should be followed by serial ultrasound examinations, and the fetus should be evaluated and treated postnatally. If moderate to severe oligohydramnios develops, the fetus should undergo complete prognostic evaluation to assess the potential for normal renal and pulmonary function at birth. If the ultrasound demonstrates presence of dysplasia, aggressive obstetrical care or prenatal decompression is not indicated. When preserved renal function is predicted, early delivery for postnatal decompression is indicated if the lungs are mature. Early delivery usually does not compromise pulmonary function as long as amniotic fluid volume has been maintained (1). If the lungs are immature, however, in-utero decompression can be considered.

**METHODS OF INTERVENTION**

**Urinary Tract**

Early attempts at bladder decompression in the late 70’s and early 80’s attempted a Seldinger-type procedure, but with limited success. A tight fitting double pigtail catheter placed over a puncture needle using a pusher worked; although it was far from ideal. Due to the difficulties in catheter placement, migration and plugging, Malecot-type and external coil type catheters were developed.

Open fetal surgery began in the early 1980’s, and was performed on eight highly selected cases of obstructive uropathy from 18-24 weeks gestation. Unfortunately this method of treatment carried significant morbidity predominantly from preterm labor (47,48,50). As a result, open fetal surgery to correct urinary tract obstruction has not since been performed. In those early patients, open decompression procedures included cutaneous vesicostomy in 7 and bilateral ureterostomies in 1. Only 4 had prolonged return of normal amniotic fluid and had adequate pulmonary function at birth. Of these only two have normal renal function at ages 5 and 8 years (51).

With advances in endoscopic equipment, the technique of transuterine endoscopy was developed at the University of California, San Francisco (52).
MacMahon and associates reported a similar fetoscopic approach in a human fetus with prune belly syndrome and oligohydramnios at 17 + weeks. They used a Neodymium-Yag laser to create a vesicoamniotic shunt, which was successful at restoration of the amniotic fluid volume. The fistula closed by 33 weeks and the child was delivered early with normally developed lungs (53). Fetal cystoscopy and valve ablation has been reported. Both antegrade and retrograde techniques have been reported. Flexible and rigid instruments were used as well. Significant fetal mortality was reported (54).

Myelomeningocele

Repair of MMC has been attempted both endoscopically and through open surgery via a hysterotomy. While technically possible, surgery for MMC is not presently being performed via a fetoscopic approach. Fetoscopy is limited by the need for multiple port sites, which can lead to membrane fixation and rupture as the uterus enlarges. In addition it is difficulty to visualize large spinal defects and requires prolonged operative times (55).

Open fetal surgery requires careful planning (Figure-2). Attempts at enhancing fetal lung maturity are made through the use of preoperative glucocorticosteroid administration to the mother. Broad spectrum antibiotics and balanced anesthesia allow the procedure to take place. The amniotic fluid is removed and kept in sterile warm syringes. A standard neurosurgical closure is performed through an approximately 8 cm hysterotomy. The neural placode is dissected from the adjacent arachnoid tissue and placed in the spinal canal. The dura is then dissected off for another layer of coverage. The skin then is freed for a final layer of closure. The amniotic fluid and added antibiotics are replaced and the uterus closed. Phophylactic tocolytics are then used (56).

Results and Complications

Urinary Tract

Early results of prenatal bladder shunting reassured physicians that the procedures could be performed safely and that catheter drainage was well tolerated in most cases. Whether prenatal shunting improves outcome remains a different matter. Patient selection is critical. One must find a dilated urinary tract with severe enough obstruction to compromise renal and pulmonary function at birth, and yet not so severe that renal function cannot be salvaged with decompression (1).

Reviews by Coplen, McLorie and Baskin have shown several things. First, obstruction and dysplasia are difficult to predict. Second, while technically feasible, fetal interventions were associated with only a 47% survival rate and a 45% of fetuses had complications (39,56-58). Third, even though oligohydramnios could be reversed, the ability to sustain good renal function was variable. Lastly, specific prenatal parameters that were effective in predicting good renal function have note been found (52,58).

The most common complication arising from open in utero fetal intervention is the instigation of
preterm labor (50). Catheters can fail either by plugging. Incorrect placement despite ultrasound guidance has resulted in fetal injury, and death (59). Reinsertions increase the risks of fetal injury and infections. Chorioamnionitis can sometimes result in pregnancy termination (51).

**Myelomeningocele**

Prenatal surgery for myelomeningocele has yielded some unexpected outcomes. Tubbs et al. were not able to show improved lower extremity function in patients that underwent intrauterine intervention (59).

Bruner et al. compared 29 fetal surgery patients with 23 controls matched for level of defect, diagnosis, calendar time and practice parameters (60). They reported a statistically significant ($P = 0.01$) decrease in the need for ventriculoperitoneal shunt placement and a lower incidence of hindbrain herniation ($P = 0.001$). Patients who underwent fetal surgery did however have a higher risk of oligohydramnios (48% vs. 4%; $P = 0.001$), and admissions for preterm contractions (50% vs. 9%; $P = 0.002$). They were also more susceptible to prematurity. Age at delivery was earlier (33 vs. 37 weeks; $P < 0.001$) and birth weight were lower (2171 vs. 3075 gm; $P < 0.001$) (56).

The effect of fetal intervention for myelomeningocele on postnatal bladder function has been studied (57,58). Despite the early repair, patterns of abnormal bladder function were exhibited. One still sees poor compliance, poor detrusor contractility, detrusor-sphincter dysynergia, hydronephrosis and vesicoureteral reflux. The previously described global defect in bladder development makes success of fetal surgery to preserve or improve bladder function unlikely. Additional studies are ongoing.

**CONCLUSIONS**

The field of fetal medicine has grown over the past two decades. Well-defined animal studies have yielded clues to the natural history and pathogenesis of obstructive uropathy and the efficacy of interventional techniques to ameliorate the sequelae of such obstruction. With advances in technology, the complexity of anomalies, which can be treated, has increased, as evidenced by the growing experience with fetal myelomeningocele repair. In addition these new scenarios provide new ethical challenges. Carrying out procedures in human fetuses must continue to be appropriately cautious and circumspect. The uncertainties and true pathologic processes surrounding urinary tract obstruction must continue to be explored. More reliable methods of determining fetal renal function lay on the horizon. Interventional techniques continue to evolve and improve. Because of the potential risks for preterm labor and maternal compromise, fetal surgery should continue to be performed only for carefully selected cases at centers that are equipped with a multidisciplinary health care team committed to ongoing, well-designed research protocols.

**REFERENCES**


38. Crombelholme TM, Harrison MR, Gobbus MS, Longaker MT, Langer JC, Callen PW, et al.: Fetal in-


