Successful treatment of lower urinary tract obstruction with peritoneal–amniotic and vesicoamniotic shunting

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ARTICLE INFO

Article history:
Received 9 January 2015
Received in revised form 13 January 2015
Accepted 15 January 2015
Available online 20 January 2015

Keywords:
LUTO
PUV
Vesicoamniotic shunt
Bladder outlet obstruction
Pigtail catheter

ABSTRACT

Lower urinary tract obstruction (LUTO), which is most commonly caused by posterior urethral valves or urethral atresia, is often associated with high perinatal mortality and morbidity and long-term childhood renal dysfunction or even renal failure. LUTO is an uncommon clinical entity that can be diagnosed by ultrasonographic examination and is particularly presented by enlarged bladder, hydronephrosis, hydrourer and oligohydramnios. Despite being a rare pathology of the genitourinary tract, detection rates of it is higher than other congenital abnormalities with ultrasonography. Moreover, ultrasonography has become the most widely used diagnostic tool in the diagnosis of LUTO. Pulmonary hypoplasia resulting from severe oligohydramnios is the most common cause of death. Bladder drainage has a critical role on perinatal survival in fetuses with LUTO especially in those predicted to have a poor prognosis. Further interventions and observations of renal therapies may be required in childhood.

1. Introduction

Genitourinary tract abnormalities are one of most detectable anomalies by sonography[1]. Lower urinary tract obstruction (LUTO) comprises a heterogeneous group of pathologies including posterior urethral valves (PUV) and urethral atresia[2]. Most causes of LUTO are PUV and urethral atresia[2]. Long-term obstructive effect of LUTO extensively induces renal parenchymal damage which, in turn, engenders to renal dysfunction and renal failure and bladder dysfunction in childhood. Oligohydramnios and pulmonary hypoplasia are two parameters of poor prognosis associated with high fetal mortality and morbidity[3]. In utero shunting is a significant antenatal intervention which has been applied for obstructive uropathy up to now[4]. Indications, ideal method of treatment and appropriate time for in utero shunting are critical points for both perinatal survival and renal and bladder development, however postnatal long-term poor renal function may render a higher risk[5]. We report two cases performed in utero shunting for LUTO.

2. Case report

2.1. Case 1

A 28-year-old woman G2P1 at 25 weeks of gestation admitted to outpatient clinic with abnormal ultrasound screening findings concerning gross fetal ascites and megacystis without oligohydramnios. Fetal termination had been offered due to perceived poor prognosis by the perinatologists in another healthcare facility. Megacystis without oligohydramnios, gross urinary ascites due to fetal bladder rupture, bilateral diaphragm elevation which caused fetal thoracic and cardiac compression and bilateral fetal pyelectasis were identified in our ultrasound screening (Figure 1). We performed amniocentesis for fetal karyotyping. After excluding major chromosomal anomalies we performed peritoneal–amniotic shunting by ultrasound–guided placement of double–ended pigtail catheter (Figure 2). Location of
catheter and fetal viability were confirmed after procedure through subsequent serial ultrasound scans. Fetal ascites, megacystis and bilateral pyelectasis resolved during the following weeks. A healthy newborn was delivered vaginally at the 38 weeks gestation. Additionally abdominal ultrasound scan of the newborn was normal.

**Figure 1.** Ultrasound image of fetal bladder rupture. 

a: Fetal ascites without oligohydramnios; b: Fetal ascites.

**Figure 2.** A transaxial scan of the fetus after peritoneal–amniotic shunting in the following weeks.

**2.2. Case 2**

A 24-year-old woman G1P0 at 19 weeks of gestation was referred from another obstetrics clinic to our outpatient clinic with abnormal ultrasound screening findings concerning fetal ascites and megacystis accompanying to oligohydramnios. Megacystis with oligohydramnios, gross fetal ascites, bilateral fetal pyelectasis were identified in our ultrasound screening (Figure 3). We performed amniocentesis for fetal karyotyping which revealed a normal chromosomal structure. Subsequently, vesicoamniotic shunting by ultrasound–guided placement of double–ended pigtail catheter (Figure 4). Fetal viability and catheter localization were ensured after the procedure, also with subsequent ultrasound scans. Fetal ascites, megacystis and bilateral pyelectasis were not identifiable during the following weeks. Healthy full–term newborn was delivered vaginally at another obstetrics clinic.

**Figure 3.** Ultrasound image of enlarged bladder with oligohydramnios.

**Figure 4.** A transaxial scan of the fetus after vesicoamniotic shunting.

**3. Discussion**

LUTO is associated with high perinatal and childhood mortality and morbidity including bladder dysfunction or renal damage[5]. Excessive accumulation of urine and high pressure arising from bladder outlet obstruction causes to deterioration of the development of bladder and renal structure, bladder dysfunction, bilateral hydronephrosis with bilateral hydroureter and subsequent renal dysfunction and renal failure[5,6]. Development of urinary organs have associations with time period and degree of obstruction[7]. In our first case, bladder rupture, gross ascites, fetal thoracic and cardiac compression due to elevation of diaphragm were identified because of long–term obstruction. The two most common causes of LUTO are PUV and urethral atresia, other less common ones are prune belly syndrome and prostatic atresia[3,8]. The PUV is more common among male than female fetuses. Our both cases were male fetuses. Prenatal ultrasound plays improvant role on diagnosis and fetal in utero intervention for LUTO. Diagnosis time of LUTO ranges from 13 to 38 weeks, but most diagnoses are established at
20–22 weeks gestation because of being time interval for routine detailed fetal anomaly scan.[2,9]. Our cases admitted to outpatient clinic at 19 and 25 weeks gestation. Diagnosis should be evaluated by karyotype analysis and aspirated urine together.[10]. For both cases we performed fetal karyotyping after amniocentesis. To evaluate renal function, serial fetal urine analysis of fetal urine is recommended inasmuch as LUTO has a progressive character.[11]. In prognosis two most substantial causes related with higher mortality rate were oligohydranrnios and pulmonary hypoplasia.[3]. Oligohydranrnios was identified in our second case. Morris et al., in their study demonstrated that all neonatal deaths were caused by pulmonary hypoplasia.[2]. Because of mortality and morbidity due to pulmonary hypoplasia and renal dysplasia by obstruction–induced pressure, the appropriate timing for in utero intervention is critical. The main choices of in utero therapies include vesicocentesis, vesicoamniotic shunt, open foetal bladder surgery and fetal cystoscopy.[5,8,9]. Double-ended pig tail catheter is the most commonly used catheter to achieve bladder drainage. Sometimes double–basket catheter can be used for long–term shunting. Won et al., demonstrated in their cases that double–basket catheter was effective for vesicoamniotic shunting for long–term decomposition.[10]. In our case, we used double–ended pig tail catheter and we did not observe and catheter complication. Vesicoamniotic shunts are most common in utero interventions to relieve the obstructive uropathy. Morris et al., showed perinatal survival is increased significantly by fetal bladder drainage (OR: 2.53).[2]. And bladder drainage versus non–treatment with an OR of 3.86 is demonstrated in systematic observational studies.[8,9]. After shunting, symptoms caused by obstruction disappeared in both cases. LUTO sometimes may be associated with certain congenital syndromes, such as prune belly syndrome, which is characterized with bladder outlet obstruction and abdominal muscle deficieny.[12]. Like our peritoneal–amniotic shunt case, Livingston et al., demonstrated bladder rupture and massive urinary ascites which caused thoracic and cardiac compression and subsequent potential hydrops fetalis after bladder outlet obstruction associated with mirror syndrome.[13]. Obstructive uropathy was relieved after peritoneal–amniotic shunt.[13]. Despite increasing perinatal survival, vesicoamniotic shunt might not have a long–term benefit.[2]. Renal dysfunction and renal failure and bladder dysfunction can develop during postnatal follow–up years.[14].

As a conclusion, the prenatal bladder drainage may improve perinatal survival in fetuses with LUTO. An appropriate timing of shunting is critical to preserve normal in utero renal development. Improving perinatal survival via shunting is clear but long–term renal functional outcomes are sceptical and poor towards increasing rates of childhood morbidity based on observational studies.

**Consent**

Written informed consent was obtained from the patient for publication of this case report and any accompanying images.

**References**


