**Case Report**

**Congenital Segmental Giant Megaureter Presenting as Cystic Abdominal Mass in a Child**

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**Abstract**

A six years old male child presented with gradually progressive distension of abdomen since one year of his age. Imaging suggested it as a large mesenteric cyst. Further evaluation and intra-operative findings suggested it to be bilateral congenital megaureter with giant one on the left side which was causing gross abdominal distension. Excision of left megaureteric segment, psoas hitch and ureteroneocystostomy was performed. Excision, tapering and ureteroneocystostomy of right ureter was performed. Congenital giant megaureter should be considered as one of differential diagnosis in children presenting as abdominal mass.

**Keywords:** Giant megaureter; Renal descensus; Ureteroneocystostomy; Psoas hitch; Abdominal mass

**Introduction**

Megaureter is a generic term indicating the presence of an enlarged ureter with or without concomitant dilatation of the upper collecting system. The normal diameter of ureter is usually 5 mm in children and rarely exceeds this size. In practice, a ureter with a diameter of 7 mm or more should be considered as a megaureter [1,2]. Megaureter may be categorized as primary or secondary, refluxing or nonrefluxing, obstructed or non-obstructed, and nonrefluxing obstructed. In all of these conditions ureteric dilatation is found completely. However, segmental megaureter or segmental cystic dilatation of ureter is a very rare entity and only a few case reports were reported in the literature [3].

**Case Report**

A six years old male child presented to us with tense grossly distended abdomen with dilated superficial abdominal veins which had been gradually progressive since infancy. He had no history of dysuria and voiding problems. His hematological and renal function tests including urine routine and urine culture sensitivity were normal. Ultrasound (USG) and Contrast Enhanced Computed Tomography (CECT) scans of abdomen (Figure 1a) showed large cystic abdominal mass suggestive of large mesenteric cyst with bilateral Hydronephrosis (HDUN) secondary to obstruction by the mass. USG guided percutaneous pigtail drainage of cystic mass was done as the child was having respiratory discomfort. Three litres of clear fluid was immediately drained and abdomen got decompressed. Drain fluid creatinine was found to be 40 mg/dl which was suggestive of urine. Repeat CECT abdomen (Figure 1b) after a week showed decompressed cystic mass, bilateral HDUN, faint contrast enhancement of left kidney and the tip of the catheter in the enlarged segment of left ureter. Nuclear scan of kidneys showed fifteen percent split function of left kidney. Micturating cysstogram (Figure 2a) showed no reflux and normal voiding pattern with no outflow obstruction. Accidentally in the meantime, the pigtail catheter got dislodged and abdomen got gradually distended again. Intraoperatoratively, the lower half of left ureter was found hugely dilated with contained urine (Figure 2b) occupying the whole of abdomen with mildly dilated upper half and a small left kidney. Right ureter was also dilated in the whole course with normal right kidney. Excision of dilated portion of left ureter, left renal descensus, psoas hitch and left ureteroneocystostomy was done. Excision and tapering and right ureteroneocystostomy was also done. He had uneventful recovery period.

On six months follow up after the surgery, the boy is doing well with normal renal function and no recurrence of distension of abdomen. CECT scan shows bilateral enhancing kidneys (Figure 3a) with excretion of contrast in bilateral ureters (Figure 3b).
Posterior Urethral Valve (PUV) or Neurogenic Bladder (NGB) that impedes ureteral emptying. Primary refluxing megaureter occurs due to a failure of the anti-reflux mechanism at the UVJ, while secondary refluxing megaureter occurs secondary to PUV and NGB when elevated bladder pressure causes incompetence of the UVJ. Megaureter may be associated with unilateral renal agenesis, complete or incomplete duplex system, ectopic kidney, contralateral cystic and dysplastic kidney, horseshoe kidney, or Hirschsprung’s disease [4]. Imaging studies are useful in the diagnosing the cases of megaureter such as ultrasound, Voiding cystourethrogram (VCUG), diuretic nuclear renography, Intravenous Pyelogram (IVP), contrast enhanced CT, and MRI. Important differential diagnoses with megaureters include mesenteric and adnexal masses in females and dilated bowel loops.
Surgical therapy is indicated for the megaureters that are secondary to severe vesicoureteric reflux or obstruction, and it is usually managed with ureteral reimplantation. Occasionally, nephro-ureterectomy may be indicated when the function of the kidney is severely impaired.

Only a few case reports have been found about segmental megaureter or segmental cystic dilatation of ureter in the literature. In 1995, a case of congenital segmental megaureter with sparing of the proximal and distal ureter, as in our case, has been reported by Ramaswamy et al. [5]. A similar case in a neonate with congenital giant megaureter with duplex kidney presenting as cystic abdominal mass has been reported by Annigery et al. [6], which was treated with left upper moiety nephroureterectomy? Huang [7] analysed congenital giant megaureter in 21 children, age ranged from 2 months to 8 years. The associated kidney or moiety was hypoplastic, dysplastic, or the site of comparatively mild hydronephrosis associated with atrophic renal parenchyma. Function and volume of the bladder was normal. Twelve of 21 patients had a duplex collecting system on the affected side. Depending on whether the giant segmental ureteral dilatation was multiple or single, these cases have presented either with marked protuberance of the mid and lower abdomen, which was distended by transverse elongated cysts, or with a round cyst mass in the lower abdomen. Nineteen patients treated by excision of the giant megaureter and the associated kidney or moiety were free of symptoms postoperatively. Several theories have been speculated regarding the pathophysiology of segmental megaureter. Ramaswamy suggested that segmental megaureter was a variant of nonrefluxing megaureter and attenuated nexuses and thin myofilaments might be responsible from this entity, as in their case. Although, in another study, it was suggested that ganglionosis might be liable the absence of ganglion cells has not subsequently been confirmed. However, there is no information about innervation patterns in dilated ureteric segment. The treatment of segmental cystic dilatation of ureter should be planned according to the ipsilateral kidney function and length of the normal ureteric segment. If the remaining parts of distal and proximal ureter have enough length, end-to-end ureteroureterostomy after excision of the segmental megaureter should be performed. If excision is not possible, tailoring or trimming of the dilated segment should be considered. If the ipsilateral ureterorenal unit is poorly functioning, hypoplastic or dysplastic, a nephroureterectomy should be planned. Segmental cystic dilatation of ureter must be distinguished from ureterocele and other conditions such as primary megaureter or ureteral diverticulum. At the same time, in the cases of segmental megaureter, the patient should be investigated for possible associated urinary system anomalies such as megacalycosis, duplication of collecting system and hypoplastic, dysplastic or nonfunctioning kidney. Congenital giant megaureter should be considered as one of differential diagnosis in children presenting as abdominal mass.

References