Multiple urinary calculi caused by primary hyperoxaluria

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Introduction

Urinary calculi are uncommon in children; accounting for 0.13-0.94 cases per 1000 hospital admissions in the western world¹. They can occur secondary to urinary tract infection, congenital urological anomalies, inborn errors of metabolism and immobilization. Commonest underlying metabolic abnormality is hypercalciuria. Cystinuria, xanthinuria, primary hyperoxaluria and hyperuricosuria are less common. Although most stones are considered to be ‘idiopathic’ in origin, it is believed, that this number would decrease if more detailed investigations of these patients are undertaken. We report two cousins with multiple, recurrent urinary calculi due to primary hyperoxaluria.

Case history 1

A nineteen month old boy from Galenbindunuwewa presented to Anuradhapura General Hospital with failure to thrive since the age of 7 months. He was born at term, weighing 2900g to healthy, first degree consanguineous parents. The weight gain was satisfactory, until 7 months of age, since when it gradually started crossing centile lines downwards, despite successful weaning, absence of significant illness or family disturbance. The stools were of normal frequency and consistency while evidence of food allergy was absent. His cousin has undergone bilateral pyelolithotomy for removal of multiple renal calculi at 4 years of age. (See case history 2).

On examination the weight, length and OFC were 7kg, 71cm and 44.5cm respectively, all below 0.4th centile. He was pale, and blood pressure was 90/60 mmHg (50th centile). Examination of other systems was normal.

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Urine microscopy revealed 100 pus cells per high power field and culture grew Escherichia coli > 10⁵ colonies. Ultrasound scan showed a single calculus measuring 5.1cm x 1.8cm in the right and multiple calculi in the left pelvi-calyceal systems. X ray KUB confirmed the calculi in the left kidney, but those on right were obscured by bowel shadows (Figure 1). IVU confirmed the presence of bilateral renal calculi.

Figure 1 X ray  KUB

Haemoglobin was 9g/dl with a normochromic normocytic anaemia in blood picture. The results of following investigations were normal; blood urea 3.5mmol/L, serum creatinine 53 micromol/L, serum sodium 130 mmol/L, potassium 4.5 mmol/L, calcium 2.6 mmol/L, phosphate 1.23 mmol/L, urine calcium excretion 25 micromol/ kg/24hrs (normal up to 100 micromol/kg/24 hrs) and urine phosphate excretion 0.4 micromol/ kg/24 hours, arterial pH 7.3 and urinary pH 6.5. Liver function tests were normal.

Microscopic examination of repeated samples of fresh centrifuged urine for calculogenic material showed an increased amount of calcium oxalate crystals, and urinary excretion of oxalate was
markedly elevated at 15.4 mmol/24 hours/1.73m² (normal value: < 0.5 mmol/ 24 hrs/ 1.73m²). For the above test urine was collected in a container acidified with hydrochloric acid and it was repeated and results were confirmed².

Urolithiasis due to primary hyperoxaluria was diagnosed and he was started on a low oxalate diet and a daily dose of 200 mg of pyridoxine. He also underwent extracorporeal shockwave lithotripsy (ESWL) at Department of Urology of National Hospital, Colombo.

On review at 4 years of age, he was asymptomatic, weighing 11kg (50th centile) and measuring 96cm (10th centile). Ultrasound scan and x-ray KUB were normal and serum creatinine was 50µmol/L. He continues to be on a low oxalate diet and pyridoxine 200mg once daily.

Case history 2

Eleven year old cousin of the above child from Dambulla, presented to Anuradhapura General Hospital with a two month history of weight loss, anorexia and limb pain. She has undergone bilateral pyelolithotomy for removal of calculi at 4 years of age at Kandy Teaching Hospital. Unfortunately she was lost for post-operative follow up.

She was born normally at term weighing 2750g to healthy consanguineous parents. On examination she weighed 20kg (<3rd centile) and measured 131cm (25th centile). She was pale and blood pressure was 100/60mm Hg. Examination of other systems was normal.

Urine microscopy and culture were normal. Haemoglobin was 5g/100ml with normocytic normochromic anaemia in blood picture. X-ray KUB showed multiple calculi in left kidney and distal ureter (Figure 2). Bilateral nephrocalcinosis was seen in ultrasound scan.

Blood urea and serum creatinine were raised being 27mmol/L and 660µmol/L. The results of following investigations were normal; serum sodium 135mmol/L, potassium 4mmol/L, calcium 2.5mmol/L and phosphate 2.09mmol/L. Urinary excretion of oxalate was elevated at 17.3mmol/ 24 hours/ 1.73m².

Chronic renal insufficiency and urolithiasis due to primary hyperoxaluria was diagnosed and she was referred to the urologist and nephrologist. She underwent ESWL at National Hospital, Colombo and end stage renal disease (ESRD) was managed conservatively. Three months later, she developed end stage renal failure and expired.

Figure 2 X-ray KUB

Discussion

The primary hyperoxalurias (PH) are rare, autosomal recessively inherited disorders of oxalate metabolism, which cause excessive endogenous oxalate production. Increased excretion of oxalate leads to saturation of calcium oxalate in urine causing crystal formation, urolithiasis and medullary nephrocalcinosis. Disease progression causes systemic oxalosis with deposition of oxalate in parenchymatous organs, bones and retina.

Two forms of PH are recognized. Type 1 (PH 1), which is relatively more common, has an annual incidence of approximately two to three patients per million and is caused by deficiency of liver-specific peroxisomal enzyme alanine-glyoxylate aminotransferase (AGA), the gene of which, AGXT has been sequenced and located on chromosome 2q 37.3. PH 1 grossly fits three clinical presentations: (a) a rare infantile form with early nephrocalcinosis and rapid kidney failure; (b) a rare late onset form with stone passage in late adulthood; (c) the most common form with recurrent urolithiasis and progressive renal failure leading to diagnosis of PH 1 in childhood or adolescence³,⁴.

Type 2 (PH 2) is rare with only 37 cases being reported and is caused by deficiency of a universal tissue enzyme D-glyceric acid dehydrogenase.
Patients with PH 2 excrete L-glyceric acid in urine in addition to oxalate. They have a less severe clinical course than PH 1 and their predominant clinical feature is urolithiasis rather than nephrocalcinosis.

While our first patient presented with failure to thrive and multiple urinary calculi the second patient had recurrent, multiple urinary calculi, nephrocalcinosis and ESRD. Both were products of first degree consanguineous parents and children themselves were first cousins. Twenty four hour urinary excretion of oxalate was markedly elevated in both. The clinical and laboratory features strongly supported a diagnosis of PH. Non-availability of facilities to assay urinary excretion of glycolate prevented us from describing the type of PH. But, as PH 2 is assumed to be rare, and to have a more benign course, clinical features of our two patients favour a diagnosis of PH 1.

Though PH first present as urolithiasis it can rapidly progress to end-stage renal failure. Hence early diagnosis and treatment is crucial to prevent ESRD. The first steps in treatment include high fluid intake, low oxalate diet and high dose of pyridoxine. Pyridoxine (vitamin B6) catalyses AGT and increases its activity. The dosage varies from 20-600 mg/day and therapeutic success can be seen when oxalate excretion decreases. Our first patient was started on this treatment and good clinical response was seen after two years.

Supportive treatment with sodium or potassium citrate and orthophosphate; which are potent inhibitors of calcium oxalate is suggested in the literature, and we are planning to start potassium citrate in our patient.

The urological treatment of children with obstructive stone disease is still being debated, and the current therapeutic strategy is minimally invasive procedures like ESWL because recurrence of calculi is a major problem. Open surgical techniques are indicated in the case of large stones. Both patients underwent ESWL and it was successful in our first case. The second patient had also undergone pyelolithotomy seven years prior to the diagnosis of PH 1.

Management of PH after onset of ESRD is conservative with haemo or peritoneal dialysis. There may be a place for liver transplantation prior to onset of ESRD as AGT is produced in the liver. Isolated kidney transplantation does not help as the disease recurs in the transplanted kidney. Combined liver - kidney transplantation has been shown to be the only way to replace both the biochemically defective organ (liver) and the pathophysiologically damaged organ (kidney) in patients with ESRD. In an ideal world, our second patient would have been offered this treatment.

The prognosis of untreated PH is poor leading to death around the age of 20 years. Hence the aim of management, especially in the developing world, should be early diagnosis and adequate treatment.

These two cases reiterate the importance of investigating children with urolithiasis, to identify the aetiology. It also highlights the fact that this is possible to a significant extent, if our limited resources are appropriately used.

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References


