Case Report

Unusual presentation of oxalate nephropathy causing acute kidney injury: A case report

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Abstract

Oxalate nephropathy due to Hyperoxaluria and elevated serum oxalate level is a well-known cause for interstitial fibrosis, and ESRD. Conditions associated with high serum Oxalate, should be considered as a possible contributing factor for a patient’s tubular injury.

Well known cause for Hyperoxaluria including enteric Hyperoxalu ria (due to gastric bypass, chronic pancreatitis, small Bowel resection, or malabsorption, as well as depletion of enteric oxalate-degrading bacteria [e.g., Oxalobacter). Other known causes of oxalate nephropathy include primary Hyperoxaluria, ethylene glycol intoxication, vitamin B6 deficiency, excessive ingestion of vitamin C or dietary substances rich in oxalic acid, aspergillosis, prolonged renal failure and various drugs (e.g., Known medications to cause Oxalate Nephropathy are: Orlistat, Praxilene, COX-2 inhibitors).

Unusual presentation with Acute Kidney Injury with incidental finding of high serum Oxalate in a patient with a known CKD stage III, recently started on Polyethelene Glycol to treat his constipation.

Case presentation

We report a case of Acute kidney injury in a 68-year-old male, presented to Ruby Memorial Hospital (Teaching hospital of West Virginia Medical School, Morgantown, WV), who required renal replacement therapy due to oxalate nephropathy, has past medical history of diabetes mellitus type 2 (DM), Hypertension, hyperlipidemia, basal cell cancer (removed from his back), chronic back pain, required Narcotic, for which he developed severe constipation and prescribe to use Polyethelen Glycol daily 6 months prior to his admission, presented to ED with generalized malaise, vomiting, and bilateral leg weakness, for 3-4 weeks duration, found with Metabolic Acidosis, and Hyperkalemia, with elevated Serum Creatinine to 7.7 mg/dl, from baseline creatinine 1.4 mg/dl 6 months prior to admission, his hemoglobin A1C was 6.4% and he had issues since 2014 controlling his potassium, he had been taking oxycodone since 2009 when he had a shoulder injury at his workplace, using oxybutynin for urge incontinence. Patient is a non-smoker, denies ingestion any kind of alcohol, or using NSAID.

Upon presentation his physical exam showed mild tachypnea with BP 124/86, presence of Retinopathy, no signs of volume depletion or overload ,Chest exam is normal with no rales or wheezing, Heart, regular, with palpable bladder, no Asterexis, but with mild hyperreflexia, with no peripheral edema.

Initial admission laboratory workup table 1 showed potassium of 6.5 mmol/L, in acute kidney injury on a Chronic Kidney Disease stage III, with a creatinine of 7.7 mg/dL k.
blood urea nitrogen of 77 mg/dL and serum bicarbonate of 10 mmol/L. An initial urine dipstick showed small blood and 5 mg/dL no Proteinuria or Hematuria, or crystals, His potassium was treated with IV calcium gluconate, insulin with dextrose 50%, as well as a sodium bicarbonate drip at 150 mL/hr. A Foley catheter was placed upon admission, yielding 700 mL of urine, over the next few days, his GI symptoms continue as he was presented, even with gentle hydration and he had a non Oliguric urine output once a Foley catheter was placed, with next 5 days, showed no improvement on his Kidney functions.

A renal ultrasound was obtained, showing normalized kidneys bilaterally with borderline increased renal echogenicity bilaterally with multiple small stones in both kidneys, with no Hydronephrosis as it shown in figure 1. Serologic workup was ordered (complements, antinuclear antibodies, myeloperoxidase antibodies, proteinase antibodies IgG, double-stranded DNA, ANCA, hepatitis B/C, as well as urine electrolytes, SPEP, and creatinine kinase), all of which were unremarkable/negative.

On day 5, after the admission A renal biopsy was performed, given the non-improvement of his kidney functions, despite resolving his Bladder Outlet Obstruction, and IV hydration, to establish diagnosis, prognosis and for future treatment planning. His kidney biopsy figure 2 showed marked interstitial fibrosis and tubular atrophy in the absence of significant glomerular disease, suggestive of chronic Tubulointerstitial injury. There is abundant calcium oxalate deposition within tubular Luminas. While some calcium oxalate deposition can be seen as a nonspecific finding in patients with acute tubular injury, abundant calcium oxalate, as in this case, raises the possibility of oxalate nephropathy, with no evidence of Diabetic Nephropathy.

Subsequent work up based on the result of his Kidney Biopsy, a 24-hour urine oxalate was collected, showing 0.41 mmol/24 hr (0.11-0.46 mmol/24 hr) and an initial plasma oxalate was 20.4 mc mol/L (<1.6 mc mol/L).

On day 6 after the admission, Hemodialysis was initiated with 5 hours sessions, using a Tunneled catheter, as indicated to clear his serum Oxalate, and to treat his moderate uremic symptoms, with daily check on plasma oxalate levels.

Genetic testing was sent to rule out primary Hyperoxaluria as a cause of his renal interstitial oxalate deposition. Genetic results showed a UMOD gene heterozygosity, with uncertain significance [6]. Per genetic report it is uncertain whether this gene had any clinical significance [7].

After patient had received 2 weeks of Dialysis He was

| Table 2: Chemistry and CBC. | Ref. Range | Actual |
|---------------------------|------------|--------|
| SODIUM                    | Range: 136 - 145 mmol/L | 133 (L) |
| POTASSIUM                 | Range: 3.5 - 5.1 mmol/L | 6.5 (HH) |
| CHLORIDE                  | Range: 96 - 111 mmol/L | 104 |
| CARBON DIOXIDE            | Range: 22 - 32 mmol/L | 13 (L) |
| BUN                       | Range: 8 - 25 mg/dL | 77 (H) |
| CREATININE                | Range: 0.62 - 1.27 mg/dL | 7.10 (H) |
| GLUCOSE                   | Range: 65 - 139 mg/dL | 127 |
| ANION GAP                 | Range: 4 - 13 mmol/L | 16 (H) |
| BUN/CREAT RATIO           | Range: 6 - 22 | 11 |
| EGFR                      | Range: > 60 mL/min/1.73m²/2 | 7 (L) |
| CALCIUM                   | Range: 8.5 - 10.2 mg/dL | 8.1 (L) |
| MAGNESIUM                 | Range: 1.6 - 2.6 mg/dL | 1.0 (LL) |
| PHOSPHORUS                | Range: 2.3 - 4.0 mg/dL | 4.7 (H) |
| WBC                       | Range: 3.7 - 11.0 x10³/uL | 7.3 |
| HGB                       | Range: 13.4 - 17.5 g/dL | 8.6 (L) |
| HCT                       | Range: 36.9 - 52.0 % | 26.6 (L) |
| PLATELET COUNT            | Range: 150 - 400 x10³/uL | 192 |

| Urinalysis | Range | Actual |
|------------|-------|--------|
| COLOR      | Normal | Normal |
| SPECIFIC GRAVITY, URINE | Range: 1.005 - 1.030 | 1.008 |
| GLUCOSE    | Negative | mg/dL |
| BILIRUBIN  | Negative | mg/dL |
| KETONES    | Negative | mg/dL |
| BLOOD      | Negative | mg/dL |
| PH URINE   | Range: 5.0 - 8.0 | 5.0 |
| PROTEIN    | Negative | mg/dL |
| UROBILINOGEN | Negative | mg/dL |
| NITRITE    | Negative | Negative |
| LEUKOCYTES | Negative | WBCs/uL |
| CREATININE, UR RAND | mg/dL | 22 |
| SODIUM, URINE RAND | mmol/L | 62 |
| OSMOLALITY, URINE | Range: 50-1,400 mOsm/kg | 218 |
ultimately discharged from the hospital with arrangement for outpatient dialysis, and he is still not declared End Stage Kidney Disease.

Discussion

In this 68-year-old male, it is unclear as to the direct cause of this patient’s elevated serum Oxalate. It is speculated that perhaps long-term supplementation with vitamin D could have increased patient’s absorption of oxalate due to increased calcium absorption [5] and deprive the small bowel from the Calcium necessary to bind to the enteral Oxalate. Polyethylene glycol, although rarely, can be depolymerized by anaerobic bacteria to ethylene glycol and diethylene glycol [8] that can be easily absorbed and converted in the liver into multiple substrates including calcium oxalate by enzymes (alcohol dehydrogenase, aldehyde dehydrogenase, lactate dehydrogenase).

In this patient’s case his Serum Creatinine was at his Baseline 1.4 mg/dl before starting his Polyethylene Glycol, and increased to above 7 mg/dl at the time of presentation, with other factor or factors could have contributed including his CKD 3, chronic Bladder outlet obstruction, with a kidney biopsy showed clear evidence of Oxalate Nephropathy, and while his high serum Oxalate was most likely multifactorial due to the patient being stage 3 CKD and being already prone to oxalate loading from his Polyethylene Glycol, in the absence of Hyperoxaluria gene.

In Chronic Kidney Disease As the glomerular filtration rate falls below 30 to 40 mL/min per 1.73 m², plasma oxalate levels increase because of reduced urinary oxalate excretion [9]. When plasma oxalate exceeds 30 micromol/L, which is the plasma supersaturating threshold for calcium oxalate, calcium oxalate is deposited into other tissues including the retina, myocardium, vessel walls, skin, bone, and the central nervous system.

It is unusual for Oxalate Nephropathy to present in advance age, so the conclusion of Multifactorial high serum oxalate level would be presumed as the trigger factors for his Acute Kidney Injury.

Long term outcome of his kidney functions, would be determine based on follow up during his outpatient Hemodialysis, after eliminating all risk factors for his high serum Oxalate, by stopping his Polyethylene Glycol, and perform Oxalate clearance of his serum Oxalate by using Hemodialysis, and treating his Bladder outlet obstruction.

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6. UMOD is a gene that has been associated with autosomal dominant UMOD-related tubulointerstitial kidney disease, previously known as familial juvenile hyperuricemia (gouty) nephropathy type 1, as well as linked to autosomal dominant glomerulocystic kidney disease with hyperuricemia and isoosmhenuria (2).
7. Autosomal Dominant Tubulointerstitial Kidney Disease, UMOD-Related Synonyms: ADTKD-UMOD, Familial Juvenile Hyperuricemic Nephropathy 1, Medullary Cystic Kidney Disease 2, UMOD-Associated Kidney Disease, Uromodulin Kidney Disease Anthony J Bleyer, MD, MS, P Suzanne Hart, PhD, and Stanislav Kmoch, PhD.
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