Gross Hematuria Post-Pieloplasty as Single Manifestation of Primary Renal Candidiasis

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Abstract
Primary Renal Candidiasis (PRC) has been described in pre-term and term newborn hospitalized in neonatal intensive care units (NICU). A risk factor for PRCs is hydronephrosis. The aim of this paper is to present the case of a toddler suffering from PRC without hospitalization in intensive care units, and with gross hematuria postpyeloplasty.

A 3 year-old male was admitted in the hospital with fever and gastroalimentary vomiting. The diagnosis of hydronephrosis by ureteropelvic junction obstruction was established. Treatment with ceftriaxone and nephrostomy guided by ultrasonography (US) was applied. Three months later, pyeloplasty and pigtail placement were carried out. Two weeks after postpyeloplasty, the patient presented gross hematuria, with secondary anemia (Hb 6.5g/dl). Urine cultures positive for Candida albicans. US renal report fungoma of 7x5mm in left kidney. Management was done with fluconazole at 6 mg/kg/day by 6 weeks. Currently asymptomatic.

Keywords: Gross hematuria; Renal candidiasis; Pyeloplasty
Abbreviations: PRC: Primary Renal Candidiasis; US: Ultrasonography; KBUS: Kidney-Bladder-Ultrasonography; APD: Anterior-Posterior Diameter; NORA: Normalised Residual Activity; DRF: Differential Renal Function

Introduction
Primary renal candidiasis (PRC) has been described previously [1]. Preterm newborn have risk factors for developing systemic or non-systemic candidiasis, because they are subject to intravenous broad spectrum antibiotics, parenteral nutrition, central venous catheters; among other factors. An important history in these patients is the presence of hydronephrosis [2].

The main clinical manifestation reported in the PRC has been anuria or oliguria secondary to mycetomas, fungomas obstructing the ureteropelvic junction, especially in pre-term patients [3,4].

The aim of this paper is to present the case of a toddler with PRC manifested by gross hematuria postpyeloplasty and being an outpatient.

Case Report
A 3 year-old male patient, no reportof prenatal hydronephrosis, with no relevant prenatal and perinatal history. Fever of 39 °C, 48 hours of evolution and gastroalimentary vomiting.

Physical examination
Palpable mass on the left flank, painful, mobile, non-stony.

Laboratories
Hematic biometry: Leukocytes 16,340/ul, neutrophils 88%. Blood chemistry: Serum creatinine 0.26 mg / dl. Uroanalysis: Leukocyte sterease (+), countless leukocytes.

Work-up
Kidney-Bladder-Ultrasonography (KBUS): Left kidney with severe hydronephrosis with pelvis of 350 ml, anterior-posterior diameter (APD) pelvic of 75mm, renal parenchyma thinned (4mm).

Diuretic renography with 99mTcMAG-3: Normalised residual activity (NORA) at 60 minutes>1 in the left kidney with 39% of differential renal function (DRF).

Based on these results, we decided to perform nephrostomy and intravenous (IV) treatment with ceftriaxone for 14 days, thus resolving the infectious condition.

Three months later, Anderson Hynes pyeloplasty and pigtail catheter placement were performed. After 24 hours postsurgery, were no complications, no incidents or accidents.
**Evolution**

After two weeks post-pyeloplasty, he presents gross hematuria with clots, without abdominal pain. During physical examination with heart rate (HR) of 125 bpm, abdomen without bleeding in surgical wound, no hematomas, no palpable mass.

**Laboratories**

Hb: 6.5 g/dl, hematocrit 19%, leukocytes 4300/ul, platelets 188,000/ul TP: 14.2, TTPa: 38.5, Urine culture: 48-hour without bacterial growth.

**Work-up**

KBUS with APDpelvic of 48 mm, no evidence of renal pelvic clots, no retroperitoneal extra-renal collections.

We decided to hospitalize with absolute rest, red blood cells were transfused; he was discharged from hospital at 48 hours without hematuria.

After 8 weeks post-pyeloplasty, a pigtail catheter was removed by cystoscopy.

After 10 weeks post-pyeloplasty, there was gross hematuria again, without clots, without symptomatic anemia.

**Laboratories**


Abdominal US to search fungomas: Two hyperechoic images in left renal parenchyma of 7x5 mm and 3x3 mm, without acoustic shadow suggestive of mycetomas (Figure 1).

**Doppler echocardiogram: Normal.**

Management with fluconazole 7 days IV and then 5 weeks oral.

Actually in the last follow-up at 12 months after the antifungal treatment, the patient was asymptomatic with negative urinalysis and urine cultures, KBUS with residual image of 3x4 mm in left kidney.

**Discussion**

PRC has been reported in term infants [5], preterm infants [6] and, infants [7], commonly in intensives care units; but we do not know of reports in a toddler as in the case that we are reporting.

Due to the use in the intensive care rooms, of broad spectrum antibiotics, intravascular devices and parenteral nutrition, among other factors, fungic systemic or localized infections have been increasing [8,9]. But our patient does not belong to this group of inpatients; he is a patient diagnosed from the outpatient clinic. Most of the reported cases of PRC in children have as a common denominator; the presence of ipsilateral hydronephrosis [10], similar to our patient; but also we added a risk factor that was the use of nephrostomy for 3 months, which undoubtedly contributed to the colonization of the kidney by candida.

PRC can manifest as pyelonephritis and/or mycetomas [11]. In children we do not know of reports of pyelonephritis (PN) as the only manifestation, but we do of PN plus mycetomas [10], or mycetomas at the ureteropelvic junction causing anuria or oliguria [12,13]. The patient in the study, did not have obstruction of the urinary tract, he had gross hematuria with acute anemia. We found reports in adults of macroscopic hematuria in PRC [14], but according to the review in the world literature this is the first case reported with this clinical manifestation in toddler.

The diagnosis of PRC has been established by ultrasonography (US), retrograde pyelography, excretory urography, anterograde pyelography and tomography [15]. The most used method is the US. We established the diagnosis by US, and also followed up with US.

The treatment of primary renal candidiasis is based on antifungal agents, the mostly used are fluconazole, amphoterin and flucytosine [16,17]; we used fluconazole for 6 weeks with complete resolution of symptom and mycetomas.

For the cases of obstructive uropathy secondary to mycetomas, several treatments have been described, such as percutaneous nephrostomy [18], percutaneous extraction of mycetomas with thrombectomy devices [19]; none of these procedures was necessary to use in our patient; gross hematuria finished with oral fluconazole treatment.

**Conclusion**

This is the first report of a child with primary renal candidiasis, manifested with anemising gross hematuria post-pyeloplasty. We present this manifestation of primary renal candidiasis to be considered in the differential diagnosis of a patient with hematuria and history of pyeloplasty with chronic use of nephrostomy.

**References**


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