ABSTRACT
The infestation by *Schistosoma haematobium* is common in African countries and West Asia. Its chronic phase is characterized by the deposition of eggs of the parasite in various tissues of the body causing inflammatory response, formation of granulomas and fibrosis. The disease often affects the urinary tract, presenting with hematuria and, in the terminal stage, renal failure by urinary obstruction and bladder squamous neoplasia. Since chronic infection can lead to significant morbidity, it is imperative that the physicians who serve this immigrant population become familiar with this disease. A case of an immigrant boy from Guinea-Bissau seen in a Nephrology appointment for monosymptomatic terminal hematuria is presented. The diagnosis of urinary schistosomiasis was confirmed by parasitological examination of urine and the pathological examination of bladder biopsies. After therapy with praziquantel, the patient became asymptomatic.

**Keywords:** Hematuria; *Schistosoma haematobium*; Case reports

INTRODUCTION
Schistosomiasis is an endemic parasitic disease in Africa, South America, Middle East, China, The Philippines and in some Caribbean islands. Approximately 200 million people in 74 countries are infected (1-2). There are three species of *Schistosoma* that are relevant for human pathology: *S. manson*, *S. japonicum* and *S. haematobium*. The clinical presentation varies depending on the infecting species and *S. haematobium* is mainly responsible for urinary tract infection. In Europe, schistosomiasis remains as a rare cause of hematuria in children(3). However, considering the close relation between Portugal and African countries (International Cooperation Agreement between Portugal and Portuguese Speaking African Countries), the differential diagnosis should always be considered in face of a child coming from that continent and presenting hematuria.

CASE REPORT
The authors presented a clinical case of a 10-year-old male child coming from Guinea-Bissau, who had been for 8 months in Portugal with the diagnosis of macroscopic hematuria and nephrotic syndrome. The family history comprised a paternal uncle with nonspecific renal disease. Since the age of 4, he referred occasional episodes of monosymptomatic terminal macroscopic hematuria and facial edema of spontaneous resolution, with use of nonspecific therapy. He presented no other relevant past history.

In the visit to the pediatric nephrologist he showed general good status, weight 35.3 Kg (P50-75), height 1.38 meters (P50-75) and blood pressure of 113/57 (complete blood count showed 12.7% eosinophilia), renal function and ionogram within normal range for
age. Urine analysis showed pH 6, density 1.017, proteins 150 mg/dL, hemoglobin +++, leukocytes 25 mg/dL, and sediments with many erythrocytes. Twelve-hour proteinuria was 28 mg/m²/hour. Uric acid, calcium, oxalate and 24-hour urinary phosphorus presented normal range for age. HIV-1 and HIV-2 serology and AgHBs were negative. The echogram showed right kidney with echogenic finding compatible with scar, bladder with echo in suspension and lobulated and vascularized wall thickness. To define the etiology, an urethrocystoscopy was performed and revealed trabeculated bladder with disperse, nodular, cotton-like and vascularized formations. Urine parasite test and bladder biopsies were positive for Schistosoma haematobium. Clinical pathology of renal biopsy showed abnormalities suggestive of segmental focal glomerulosclerosis.

After beginning therapy with praziquantel (40 mg/kg) single dose there was resolution of hematuria. About six months later, he remained asymptomatic with normal urinary results and 12-hour proteinuria of 3.4 mg/m²/h.

**DISCUSSION**

Urinary schistosomiasis is a serious public health problem in tropical countries and it is particularly common in Sub-Saharan Africa, resulting in high morbidity, especially in situations of chronic infection. It is estimated that about 200 million people are infected, out of which 88 million are aged less than 15 years (4). The peak of incidence and prevalence occurs in school-aged children, between 8 and 12 years (5). Male gender is the most affected owing to greater recreational exposure to water (5). Schistosoma haematobium is responsible for the infestation of the urinary tract leading to fibrosis, stenosis and calcification (1). The life cycle of this parasite is complex and includes sexual reproduction of adult parasites in humans, in addition to a stage of asexual reproduction in the intermediate host, the fresh water snail Bulinus (6). Infection in humans is acquired by direct contact with water that contains free larva forms (cercaria) released by the infected snails. Considering the preference for the venous plexus of the urinary tract, adult worms of S. haematobium live and lay their eggs there. The pathogenesis of the disease caused by this parasite is normally related with immune reactions of the body against the presence of the eggs in the tissues, which induce granulomatous inflammatory reactions (7). Adult worms are recovered by host cells (many blood groups, major histocompatibility complex molecules, immunoglobulins and albumin), which mask their own antigens, escaping from the action of the immune system and they may continue to produce an incredible number of eggs for many years (8). About 10 to 12 weeks after the contact with the parasite, there is terminal or total hematuria, which may be accompanied by dysuria, pollakiuria, or fever (1). Late manifestations (chronic schistosomiasis), in addition to hematuria, include proteinuria (many times as nephrotic manifestation), calcifications, renal cramps, hydronephrosis, urethral obstruction, renal failure and, possibly, bladder neoplasm (4). Less frequently, there is the formation of immune complexes that contain specific parasite antigens that when deposited on the glomerular capillaries induce schistosomic glomerulopathy of variable degree. This renal impairment is frequent in infestation by Schistosoma mansoni and the most severe forms of the disease (glomerulonephritis grades III and IV) are normally associated with hepatic-splenic involvement and progress to chronic renal failure (9).

The definite diagnosis of urinary schistosomiasis is made by identification of eggs in the urine or bladder biopsies. Considering that egg elimination is constant during the whole day, it is recommended the collection of three urine samples between 10 am and 2 pm (greater excretion period) or after physical exercise (3). Serological studies for schistosoma are equally useful in the diagnosis, but they do not differentiate previous from recent infection. Renal and vesical echographies are non-invasive tests that enable detection of advanced disease. In an initial stage, cystoscopy normally reveals granuloma and mucosa congestion, and later it shows sandy patches (rugous areas of vesical mucosa involving the egg deposits) (10).

In the chronic stages, egg elimination in the urine is significantly lower and it may not be present, and the diagnosis is made based on radiological tests and clinical pathology of biopsied lesions (6). Radioigraphy of the urinary tract shows calcifications at urethral and vesical level resulting from egg laying on the mucosa (10). Excretory urography aims to identify possible complications in advanced stages, such as distal urethral stenosis and superior urinary tract dilation (10).

The treatment of choice is praziquantel, a single dose of 40 mg/kg (6). This therapy provides 80% cure rate and substantial reduction of number of parasites and egg excretion when cure is not effective (1,4). The treatment with this antihelminthic eradicates the parasite and terminates the inflammatory response responsible for the chronic manifestations of the disease (5).

In the reported case, the patient had monosymptomatic macroscopic hematuria with moderate proteinuria and previous episodes of generalized poorly characterized edema. Despite the fact that the investigation of Schistosoma eggs in the urine was positive, since a more complicated disease was suspected of, it was decided to use cystoscopy and renal biopsy as well. The histology abnormalities
A rare case of hematuria

found – segmental focal glomerulosclerosis – are not normally associated with infestation by *Schistosoma haematobium* and they do not seem to respond to antiparasitic or immunosuppressant therapy, such as in this case. Moreover, the patient did not present liver or renal disease, which is normally associated with this type of glomerulopathy. However, the clinical resolution of the edema and the elimination of proteinuria, with no other therapy than praziquantel, seemed to confirm the diagnosis of schistosoma glomerulopathy. Considering the favorable progression (maintained after 6 months), the renal biopsy was not repeated.

The report of this clinical case aimed to build awareness of healthcare professionals who work with children coming from the African continent because it is a rare cause of hematuria in Portugal, but if it is diagnosed and treated immediately, chronic manifestations may be prevented, reducing urinary symptoms that lead to high morbidity and mortality, especially severe complications, such as renal failure and bladder neoplasm.

REFERENCES