Intravascular haemolysis and renal failure

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To the Editor,

It is well known that acute haemolysis is a cause of acute renal failure due to tubular damage caused by pigments being deposited in the proximal tubule. Maintained haemolysis can produce chronic renal damage, caused by different mechanisms.

We present two patients with intravascular haemolysis produced by different causes, both with acute renal failure but different progression.

A 57-year-old male, crop-sprayer, was admitted to the haematology department due to non-immune haemolytic anaemia (negative direct Coombs test). He had organophosphate poisoning, which had lasted 4 days, with oligoanuria and acute renal failure. Haemolysis stopped after two plasmapheresis sessions. Upon admission he presented with: haemoglobin (Hb): 6.3g/dl, haematocrit (Ht): 19%, leukocytes 30 630/µl (Ne 77.7%), platelets: 217 000/µl. Blood smear: intense anisocytosis, polychromatophilia, microspherocytes (7-8/field). occasional basophilic stippling, presence of erythroblasts. There were no schistocytes. Haptoglobin: <7.56mg/dl. Total bilirubin: 4.80mg/dl; direct bilirubin: 0.6mg/dl; LDH: 10 500IIU/l (Figure 1); normal iron profile; urea: 83mg/dl; Cr: 2.46mg/dl (Figure 2). Urine: proteins: 150mg/dl; haemoglobin: ++++, 47 red blood cells/field. Renal function was maintained stable with conservative treatment. Following the surgical closure, the leak stopped the haemolysis and the renal function recovered up to a glomerular filtration rate of 57ml/min, with Cr 1.7mg/dl.

Intravascular haemolysis of any cause can produce acute tubular necrosis, due to haemoglobinuria. It presents with red/brown urine and plasma, low haptoglobin, elevated LDH, deteriorated renal function and fractional excretion of sodium less than 1%. The incidence is unknown, reaching 50% in massive haemolysis.1,2

Haemoglobin is released to the plasma, binds to haptoglobin and is degraded by the reticuloendothelial system. When the haptoglobin is saturated, the free haemoglobin goes from its usual closure and chronic haemolysis. He had baseline Hb: 10.6g/dl and LDH: 1500-2000IIU/l. Baseline renal function: 71.86ml/min, with Cr: 1.85mg/dl. During this episode he presented with Hb: 7.6g/dl; haematocrit: 25.2% ; leukocytes : 7070/µl (Ne: 77.7%) ; platelets: 217 000/µl. Blood smear: intense anisocytosis, polychromatophilia, microspherocytes (7-8/field). occasional basophilic stippling, presence of erythroblasts. There were no schistocytes. Haptoglobin: <7.56mg/dl. Total bilirubin: 4.80mg/dl; direct bilirubin: 0.6mg/dl; LDH: 10 500IIU/l (Figure 1); normal iron profile; urea: 83mg/dl; Cr: 2.46mg/dl (Figure 2). Urine: proteins: 150mg/dl; haemoglobin: ++++, 47 red blood cells/field. Renal function was maintained stable with conservative treatment. Following the surgical closure, the leak stopped the haemolysis and the renal function recovered up to a glomerular filtration rate of 57ml/min, with Cr 1.7mg/dl.

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tetrameric form to a dimeric form. It filters through the glomerulus and goes inside the proximal tubule when it binds to the apical surface of megalin-cubulin receptor. It is there that globin and haem group are dissociated. The intracellular increase of proteins in the haem group produces nephrotoxicity caused by renal hypoperfusion, direct cytotoxicity and formation of intratubular casts interacting with Tamm–Horsfall protein, which obstruct the tubules.

During massive haemolysis, deleterious effects of the nitric oxide depletion are observed: smooth muscle tone imbalance, vascular constriction, thrombosis and intrarenal vasoconstriction.

Chronic damage is produced because the kidney is continually exposed to the haem group, mediated by monocyte chemoattractant protein-1 (MCP-1) and TGFβ1, which recruit monocyte and macrophages and provoke fibrosis.

These cases show the two forms kidney damage expression in haemolysis: acute and chronic. The first had acute haemolysis and required haemodialysis, with complete recovery of the renal function. The second had chronic haemolysis and chronic renal damage due to maintained exposure to the haem group, and needed conservative treatment, maintaining certain renal failure.

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Figure 2. Comparing evolution of creatinine figures in two patients.

Atypical localisation of tuberculosis in kidney transplants
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To the Editor,
Kidney transplant recipients have a greater risk of developing tuberculosis, commonly being atypical and extrapulmonary. We present the case of two patients submitted to kidney transplant with extrapulmonary tuberculosis in an uncommon localisation.

A 66-year-old female, with chronic kidney failure secondary to hepatorenal polycystosis, which received a deceased-donor kidney transplant and treatment with basiliximab, steroids, mycophenolate mofetil and tacrolimus. She suffered a type IIb cortical-resistant acute rejection and needed treatment with OKT3. After four months she was admitted for fever, general discomfort and intense asthenia. She was diagnosed with suspected pulmonary tuberculosis by chest computed tomography (CT) and fibrobronchoscopy, confirmed by Ziehl–Neelsen staining and Löwenstein culture. She was prescribed treatment with rifampicin, isoniazid and pyrazinamide for two months, followed by rifampicin and isoniazid for four months. After 15 days she was readmitted for confusion, occipital cephalgia and visual alterations. In a brain resonance, multiple hypertensive nodules were seen in T2, with nodular focal contrast in the right frontal, subcortical, suprasylvian, right occipital areas and in cerebellar peduncles, indicative of granulomatous infiltration secondary to tuberculosis (Figure 1). Treatment with isoniazid and rifampicin was extended to nine months and the patient recovered.

A 41-year-old male, diagnosed with hepatorenal polycystosis received a deceased-donor kidney transplant with immunosuppression with cyclosporin and steroids. He suffered a grade IIb acute interstitial rejection, treated with