Acute Kidney Injury in a Child with Paroxysmal Cold Haemoglobinuria

Abstract

Paroxysmal cold haemoglobinuria (PCH), first described in 1854, is characterized by acute intravascular haemolysis mediated by the Donath-Landsteiner (D-L) antibody. It accounts for up to forty percent of all cases of autoimmune haemolytic anaemia in children. Acute kidney injury (AKI) in association with PCH, however, is very rare with only two published case reports in the paediatric literature.

We report the case of a nine-year-old boy who developed AKI secondary to PCH. Supportive treatment with active warming and strict fluid management was commenced. He required two sessions of haemodialysis and two warmed red cell transfusions. Recovery was complete with normal renal profile and haemoglobin at five weeks. At twenty months he remains disease-free.

This case adds to the paediatric literature describing AKI in the setting of PCH. Although AKI is a rare complication, the importance of careful monitoring of fluid status and renal function is emphasised in the setting of acute intravascular haemolysis.

Introduction

Paroxysmal cold haemoglobinuria (PCH), first described in 1854, is characterized by acute intravascular haemolysis, mediated by the Donath-Landsteiner antibody. Autoantibody activation against the p-antigen on red cell membranes activates complement cascade, resulting in red cell membrane perforation and intravascular haemolysis [1]. Although rare, it accounts for upwards of 40% of all cases of autoimmune haemolytic anaemia in children [2]. Patients commonly present with haemoglobinuria, jaundice and anaemia after exposure to cold. Urticarial symptoms may occur as the p-antigen has been found on lymphocytes and skin fibroblasts [1]. Acute kidney injury (AKI) in association with PCH, however, is very rare with only two published case reports in the paediatric literature [3,4]. We report the case of a 9-year-old boy who developed acute kidney injury secondary to PCH.

Case History

A previously healthy 9-year-old boy presented with acute onset of lip-swelling and passing dark urine, after eating an ice-lollipop. He had a preceding upper respiratory tract infection 2 weeks prior to presentation. On examination he had evidence of angioedema, was normotensive with no generalised oedema. Initial investigations revealed a normal haemoglobin level and normal renal function. Urinalysis was positive for 2+ blood. Urinary direct microscopy, however, did not reveal any red blood cells, casts or organisms.

On day 2 he deteriorated clinically with vomiting and abdominal pain. He developed jaundice, hypertension and became oliguric. Serial investigations revealed progressive acute kidney injury with intravascular haemolysis. Serum creatinine increased from a baseline level of 35 μmol/L to a level of 233 μmol/L over 24 hours. Ultrasound renal tracts identified bilateral echo bright kidneys. The patient developed progressive anaemia, thrombocytopaenia, elevated LDH and bilirubin. He was managed with careful replacement of fluids, strict fluid balance and antihypertensive treatment.

Further haematological investigations confirmed a positive direct Coombs test for anti-C3d and haemoglobinuria. The diagnosis of PCH was confirmed on day 4 of admission. Initially no abnormalities were detected on blood film, but subsequent examination by the paediatric haematology laboratory revealed the unusual phenomenon of erythrophagocytosis (Figure 1).

This consumption of red cells by macrophages and other phagocytes is a rare in vivo phenomenon, pathognomonic of PCH [5]. It was not possible to perform the Donath-Landsteiner test in the laboratory, the gold standard diagnostic test [1], due to difficulties in transporting blood at 5 °C and 37 °C. Extensive immunology testing was otherwise normal. Creatinine kinase was normal. Rhinovirus was detected on respiratory secretions, however all other virology and

Figure 1: Blood film demonstrating erythrophagocytosis.
with haemoglobinuria, jaundice and anaemia after exposure to cold, in addition to urticarial symptoms related to p-antigen presence on skin fibroblasts [1,2]. The preceding viral illness in this case may have been rhinovirus which was detected in respiratory secretions. In particular tests for Influenza A+B, Mycoplasma, Measles, EBV, Parvovirus, Adenovirus and Coronavirus were all negative. Cold exposure from ingestion of the ice-lollipop has triggered the antibody activation leading to local angioedema and haemoglobinuria. Importantly, the latter was misdiagnosed as haematuria until urine microscopy was negative for red blood cells and casts on repeat testing.

When PCH is clinically suspected, a diagnosis is made by performing the Donath-Landsteiner antibody test, an In vitro assay for biphasic intravascular haemolysis [1]. Unfortunately our laboratory did not have the appropriate facilities to perform this investigation. It was considered unnecessary, however, as the erythrophagocytosis in association with the haemoglobinuria and positive anti-C3d on the direct Coombs test was considered diagnostic of PCH [2,5]. We emphasise the role of the paediatric haematology scientific officers in recognizing abnormalities of blood film which was misreported as normal on repeat tests in the haematology laboratory.

Treatment of PCH revolves around supportive care, avoidance of cold exposure and early recognition and treatment of complications. Warmed packed red cell transfusions should be administered for life threatening haemolysis and symptomatic anaemia [3], as was necessary for our patient. AKI is a very rare complication, however our patient required two sessions of haemodialysis in the management of severe uraemia, with subsequent gradual improvement in renal function. There is some evidence to support the use of plasma exchange therapy with 5% albumin in PCH whilst steroids, although frequently used, have not been shown to reduce the clinical course.

Complications associated with PCH include urticarial eruptions from skin fibroblast activation, severe anaemia following acute intravascular haemolysis, AKI and multiorgan failure [1]. Mortality is rare and is often secondary to multigorgan failure from severe anaemia [1]. Acute kidney injury in the setting of PCH is very rare with only 2 previous case reports in the paediatric literature [3,4]. The pathological processes by which acute kidney injury occurs in PCH and other forms of autoimmune haemolysis are poorly understood. Acute kidney injury, however, must be recognized as an important complication of PCH and other causes of acute intravascular haemolysis. It therefore follows that these patients should have strict fluid balance and renal function monitoring throughout the period of their illness.

The long term prognosis following PCH is excellent. It is thought that antibodies may persist in this condition for up to three months and hence cold avoidance is important during this period. The patient described in this case recovered well and remains disease free at 20 months since diagnosis.

This case adds to the paediatric literature describing the rare occurrence of AKI secondary to intravascular haemolysis caused by PCH. Timely recognition of PCH allows prompt, supportive therapy. We encourage the support of paediatric haematology scientific officers in diagnostic difficulties. The importance of strict monitoring of fluid balance & renal function in acute intravascular haemolysisis...
References

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