Recurrent Transient Loss of Consciousness Induced by Acute Massive Gastrointestinal Hemorrhage during Hemodialysis: A Case Report

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Key Words
Transient loss of consciousness · End-stage renal disease · Massive gastrointestinal hemorrhage

Abstract
A 67-year-old man with chronic hemodialysis (HD), who had end-stage renal disease secondary to diabetic nephropathy and had been on HD for 9 years with ultrapure dialysis solution, is presented with recurrent transient loss of consciousness during HD.

Case Report
On May 8, 2010, in the morning, a 67-year-old man on chronic hemodialysis (HD) with predialysis blood pressure 145/85 mm Hg and heart rate 62 beats/min presented with convulsion and unconsciousness 30 min after starting HD. He had end-stage renal disease (ESRD) secondary to diabetic nephropathy and had been on HD for 9 years with an ultrapure dialysis solution schedule of 3 sessions per week. He had no hemorrhagic tendency and no peptic ulcer. A routine heparin, constant-infusion method was used during HD. He was given an initial 15-mg heparin bolus dose followed by 7.5 mg/h infusion for 3 h. He had diabetes mellitus for 25 years and hypertension for 9 years. His usual predialysis blood pressure was 140–160/80–90 mm Hg. On May 6, 2010, he had undergone a conventional blood analysis, which is done once monthly in chronic HD patients. Laboratory values showed: Hb 106 g/l, Hct 32.7%, WBC 7.52×10^9/l, PLTs 297×10^9/l, serum creatinine 1,037.1 μmol/l, serum BUN 23.65 mmol/l, and serum potassium 4.9 mmol/l. An initial physical examination revealed that he suffered from obnubilation. However, pathological signs of the nervous system were negative and both of his pupils were isocoric. His vital signs were: blood pressure 92/50 mm Hg, respiratory rate 20 breaths/min, and heart rate 100 beats/min. His blood glucose concentration was 5.8 mmol/l by glucometer. The ultrafiltration rate was immediately reduced to zero and a bolus of 0.9% saline (100 ml) was rapidly administered. After the above-mentioned treatment, his blood pressure returned to 105/60 mm Hg and consciousness returned to normal. However, he then presented with another conscious disturbance after HD ultrafiltration was resumed. HD was stopped and he was transferred to our emergency department. His consciousness returned to normal again after HD ultrafiltration was stopped and a bolus of 0.9% saline (500 ml). A computed tomographic (CT) scan of his head revealed normal findings. An electrocardiogram and head ultrasound also revealed normal findings. His serum LDH, haptoglobins, and red cell fragments were all normal, but serum potassium and BUN were 7.8 and 36.07 mmol/l, respectively which were higher than 2 days earlier. He was therefore treated with urgent heparin-free HD again (in order to avoid intracerebral hemorrhage) on that same day in the afternoon. He presented with convulsion and...
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unconsciousness again 60 min after HD was initiated. HD was stopped and he was transferred to our emergency department again, where his consciousness returned to normal after a massive fluid infusion.

His blood cell count was not remeasured until the morning of May 10, 2010, when he was observed to have anemia during the next session of HD. During the same morning, his blood cell count revealed severe anemia: RBC $1.07 \times 10^{12}$/l, Hb 33 g/l, Hct 10.7%, WBC $24.61 \times 10^9$/l, and PLTs $220 \times 10^9$/l. After careful questioning, he complained of large unshaped dark stools on the morning of May 8, 2010, without abdominal pain, and after that, did not have any stools until May 16, 2010. His abdomen was flat and soft, with no tenderness. His liver and spleen were not palpable. On May 16, 2010, the routine stool test showed that occult blood was positive and there was no RBC in his stools.

Discussion

A 67-year-old man with chronic HD who presented with recurrent transient loss of consciousness during HD is reported. The most common causes of acute cerebral dysfunction during or immediately after HD are disequilibrium syndrome and intracranial bleeding [1, 2]. Because the possibility of disequilibrium syndrome is seldom seen in patients on regular HD for 9 years, the cause of his recurrent transient loss of consciousness during HD was simply and incorrectly diagnosed as an acute cerebrovascular accident. It was not until the result of his head CT scan revealed normal findings that the possibility of an acute cerebrovascular accident was excluded. An ultra-pure dialysis solution was used and the hemodialyzers were not reused in our unit. Only this patient presented a drop in Hct on that particular day, and his serum LDH, haptoglobins, and RBC fragments, which were examined in the emergency room, were all normal. Also, the rapidity of hyperkalemia was caused by decomposing and reabsorption of blood in the gastrointestinal tract, so it was not a possible explanation that his drop in Hct was caused by hemolysis. The real cause of his recurrent transient loss of consciousness during HD was hypovolemia due to an acute massive gastrointestinal hemorrhage. The attending emergency room doctor was misled by the patient’s nor-

Fig. 1. Gastroscopic morphological change of multiple superficial ulcers in the antrum.
Fig. 2. Gastroscopic morphological change of mucous hyperemia and mucosal edema around the ulcers in the antrum.
Gastrointestinal bleeding is occurring with a greater frequency and is associated with a higher mortality in uremic than non-uremic patients [3, 4]. In the US Renal Data System study of renal transplant recipients, the age-adjusted risk ratio (RR) for hospitalization due to gastrointestinal hemorrhage accounts for 3–7% of all deaths in patients with ESRD [5]. A prospective study by Fiaccadori et al. [6] found that acute gastrointestinal hemorrhage in patients with impaired renal function was associated with an increase in mortality (RR = 2.57) and a 37% increase in length of hospital stay as compared to non-renal failure patients. In patients with ESRD, the most common causes of upper gastrointestinal bleeding are peptic ulcers, hemorrhagic esophagitis, gastritis, duodenitis and gastric telangiectasias [7, 8]. This patient did not have any gastrointestinal symptoms, except gastric ulceration. Endoscopy confirmed the presence of a significant mucosal injury. There was no personal or family history of ulcer disease. There were, however, some clues with respect to the gastrointestinal hemorrhage in this patient, such as high blood urea nitrogen and hyperpotassemia even after dialysis because of toxin resorption after gastrointestinal hemorrhage.

On the other hand, renal anemia is very common in maintenance HD patients as a result of the insufficient synthesis of EPO [9]. However, use of rHuEPO increases iron demands. It is demonstrated that in more than half of the patients on rHuEPO therapy, erythropoiesis will be affected by iron deficiency [10]. Although iron storage may be normal or even increased in some patients, many lack available iron since the iron stored cannot be effectively released to meet the demands of the bone marrow for hemopoiesis. This may cause functional iron deficiency and may also reduce the efficacy of anemia therapies [11]. According to guidelines [12], all anemic, iron-deficient HD patients should be administered iron. After oral iron, all patients have dark stools and routine stool tests show that occult blood is positive without gastrointestinal bleeding because of the effect of oral iron. The use of intravenous iron is therefore recommended in maintenance HD patients with an iron deficiency to avoid the diagnosis of gastrointestinal hemorrhage.

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