335 mmol/L, respectively. Computed tomography (CT) showed diffuse cerebral swelling consistent with hemorrhagic contusion and epidural hematoma at right temporoparietal area (Fig. 1). 150 mL of mannitol solution (20%) every 4 hours was prescribed. However, despite treatment to lower intracranial pressure (ICP), that is, mannitol, mild hyperventilation, and extraventricular drainage (EVD), his mental status remained semicomatose and recurrent convulsive generalized tonic-clonic seizures developed, and thus, he was admitted to the neurosurgical intensive care unit for adequate hydration, anticonvulsant and correction

INTRODUCTION

Rhabdomyolysis causes muscle breakdown and the leakage of intracellular muscle constituents (e.g., enzymes, electrolytes, myoglobin) into the circulation and extracellular fluid\(^4\). When massive amounts of myoglobin are released, the binding capacity of plasma proteins is exceeded and the myoglobin reaches renal tubules and can cause acute renal failure\(^5\). Prolonged mental deterioration and convulsive seizures should be included among the neurological conditions in which rhabdomyolysis can develop\(^6\). A case is presented in which prolonged mental deterioration with hyperosmolar state and continuous convulsive seizure after head injury led to rhabdomyolysis and acute renal failure. The pathophysiological mechanism of this uncommon entity after severe head injury is discussed and a review of relevant literature is included.

CASE REPORT

A previously healthy 42-year-old man was referred to our emergency department due to a deteriorating mental status after a car accident. Emergency services reported decreased mental status and a Glasgow Coma Scale score of 7. There were no associated systemic injuries, muscle contusions, internal organ damage or swelling of the extremities. However, he showed decerebrated rigidity at both arms and legs. On admission, sodium, potassium, and osmolarity were 143 meq/L, 3.5 meq/L, and 335 mmol/L, respectively. Computed tomography (CT) showed diffuse cerebral swelling consistent with hemorrhagic contusion and epidural hematoma at right temporoparietal area (Fig. 1). 150 mL of mannitol solution (20%) every 4 hours was prescribed. However, despite treatment to lower intracranial pressure (ICP), that is, mannitol, mild hyperventilation, and extraventricular drainage (EVD), his mental status remained semicomatose and recurrent convulsive generalized tonic-clonic seizures developed, and thus, he was admitted to the neurosurgical intensive care unit for adequate hydration, anticonvulsant and correction

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**Case Report**

**Fatal Rhabdomyolysis in a Patient with Head Injury**

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Rhabdomyolysis is a rare but potentially life-threatening disorder caused by the release of injured skeletal muscle components into the circulation. The authors report a case of severe head injury, in which a hyperosmolar state and continuous seizure complicated by severe rhabdomyolysis and acute renal failure evolved during the course of treatment resulted in a fatal outcome despite intensive supportive treatment. Our bitter experience suggests that rhabdomyolysis should be born in mind in patients with severe head injury who may develop hyperosmolar state and continuous seizure.

**Key Words:** Rhabdomyolysis · Head injury.
of the metabolic derangement. Unfortunately, myoglobinuria developed 3 days after admission with a high serum creatine kinase (CK) level up to 48, 837 IU/L (normal 55-215 IU/L). At this time, his serum sodium, potassium, and osmolality were 152 meq/L, 3.0 meq/L and 352 mmol/L. Acute renal failure developed 3 days after admission despite intensive fluid replacement and diuresis. CK level peaked at 52, 825 IU/L on day 5, and serum myoglobin exceeded 25 mg/dL (normally <2.5 mg/dL). Despite intensive supportive treatment, the patient became anuric 5 days after admission and required hemodialysis. No source of muscle breakdown was evident, but his condition deteriorated rapidly, culminating in cardiorespiratory arrest with dilated non-reactive pupils. He died 8 days after admission.

**DISCUSSION**

Rhabdomyolysis is a frequent concomitant of multisystem trauma, and the source of muscle breakdown is often apparent from the pattern of injury. Although mechanical damage to muscle is the most common cause of rhabdomyolysis in trauma patients, other causes should not be overlooked. In patients with severe head injury, a prolonged deteriorated mental status, convulsive seizure, and hyperosmolar states importantly predispose significant rhabdomyolysis. Our patient had a decreased mental status, which probably played a role in the development of rhabdomyolysis. Experiments with human volunteers have demonstrated that sustained lying in a fixed and rigid position can cause sufficient intramuscular pressure to occlude capillaries. Furthermore, mannitol-induced hyperosmolar states have also been known to cause significant rhabdomyolysis. Our patient had received mannitol solution (20%) intravenously 150 mL/4 hr for 5 days, which probably aggravated his hyperosmolar state. Singh et al. reported relations between the hyperosmolar state and the pathogenesis of rhabdomyolysis, and observed that serum sodium and serum osmolality were both linearly related to serum CK. It was also noted that hypokalemia and hypophosphatemia were present in patients who developed rhabdomyolysis. In our patient, a hyperosmolar state, hypernatremia, and hypokalemia, were all present, and serum CK reached an extremely high level. It is known that the combination of potential potassium and phosphate deficiency in combination with acidosis, hypernatremia, and a hyperosmolar state, can reduce sodium release from muscle cells, and thus, decrease transmembrane potentials, which animal experiments have shown to be a crucial precursor of rhabdomyolysis. In addition, our patient experienced convulsive seizures and intense rigidity, and in our opinion, these factors are also likely to predispose rhabdomyolysis. Partial and generalized seizures affect autonomic function during seizures. The cause of muscle injury in exertional rhabdomyolysis has been postulated to be energy deficiency. During maximal exertion, energy is obtained almost solely from muscle glycogen, and if exertional requirements exceed available energy stores, muscle glycogen is depleted. Furthermore, if energy production is deficient, energy rich phosphate compounds are subject to subcellular competition and insufficient energy is available for muscle cell membrane structural maintenance, which results in changes ranging from reversible permeability to cell death. This type of cellular damage results in the release of intracellular potassium, purine, creatinine, myoglobin, and creatine kinase. Fever, malaise, weakness, and extremity pain, as well as swelling and tenderness are usually reported when the onset of rhabdomyolysis results from exertion. In patients with a deteriorated mental status, fever and discoloration of urine should suggest the condition.

Management of rhabdomyolysis and myoglobinuria begins with removing the cause, if possible. However, volume contraction, hypernatremia and the hyperosmolar state in neurosurgical patients treated with osmotic diuretic agents for brain swelling should be done with caution. The most appropriate treatment for hyperosmolar state-induced rhabdomyolysis is adequate hydration and alkaline diuresis. In patients with convulsive seizures or rigidity, sustained muscle contractions may be modified by the use of an anticonvulsant or muscle relaxant. Dialysis may be required if renal failure persists; indications for dialysis include hyperkalemia, uremic pericarditis, or encephalopathy, and evidence of volume overload.

**CONCLUSION**

We report a fatal case of rhabdomyolysis that developed shortly after severe head injury. Although rare, the possibility of rhabdomyolysis should be kept in mind in patients with severe head injury.

**References**