Case Report

Lose 500,000 Creatine Phosphokinase With a 2-Hour Workout: A Case Report on Rhabdomyolysis in a Novice Athlete

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Abstract

The co-occurrence of acute kidney injury secondary to rhabdomyolysis in a young patient can present as a prognostic and therapeutic challenge. Here we report a case with an unusually high creatinine phosphokinase of over 500,000 U/L after a 2-hour workout session at the gym. The challenge was to assess if the need for renal replacement therapy was warranted and whether IV fluid therapy with close monitoring would be enough to treat him, given he had a good urine output and no acid base disorder. The literature is reviewed as an attempt to delineate a rational approach to evaluating novice athletes at risk for rhabdomyolysis.

Keywords: Rhabdomyolysis; Acute kidney injury; Myoglobin; Creatinine kinase

Introduction

Rhabdomyolysis (RB) is a condition that induces a rapid breakdown of skeletal muscle cells, and involves leakage of large quantities of potentially toxic intracellular contents into plasma. It is characterized by marked elevation of creatine phosphokinase (CK), myoglobinuria, and aldolase in blood [1]. The most common symptoms are muscle weakness, myalgia and high colored urine [2]. However, up to 50% of the cases may present as asymptomatic [1]. There is no established serum CK level cut-off for diagnosing RB, but a CK level > 5,000 IU/L indicates serious muscle injury [3].

We report a case of an active young male patient who presented to Staten Island University Hospital with severe fatigue and reddish brown urine following 2 h of intense workout with a CK of over 500,000 IU/L.

Pathophysiology

During muscle injury, there is a decreased energy production that impedes the energy-dependent ion pumps, such as Na\textsuperscript{+}/K\textsuperscript{+} ATPase and Ca\textsuperscript{2+} ATPase in the sarcolemma [4, 5]. This results in an increase cellular permeability to sodium ions [5, 6]. To counter the sodium influx, the cell increases the activity of the 2Na\textsuperscript{+}/Ca\textsuperscript{2+} exchanger, which removes the excess intracellular sodium ions from the cytoplasm by exchanging it with calcium ions, eventually increasing the cytoplasmic calcium [6, 7]. Additionally, as the concentration of extracellular ionized calcium is 10,000 times higher than that in the intracellular space [8], any interruption of the plasma membrane continuity results in massive influx of extracellular calcium into the cytoplasm driven by the gradient [9].

Due to the cell’s incompetent energy-dependent ion pumps and the increased permeability to calcium, the muscle cells get overloaded with calcium ions. Calcium activates a variety of proteolytic enzymes that injure the cell membrane, allowing efflux of cellular components such as potassium, myoglobin, phosphate and CK into the circulation. Additionally, the extracellular calcium is forced to the intracellular space leading to interaction with actin and myosin causing muscle destruction [10-12].

Myoglobin is composed of globin, and a molecule of heme that is found in the muscle tissue of vertebrates, giving its characteristic color. Physiologically, the excretion of myoglobin occurs mainly through the renal system. During RB, excess myoglobin from skeletal muscle cells enters the plasma known as myoglobinemia [7]. When the concentration of myoglobin exceeds the reabsorbing capacity of glomerular cells, an excessive amount of myoglobin appears in the urine as myoglobinuria that may be detected either with a urine dipstick (microscopic myoglobinuria) or as reddish-brown urine in the case of severe RB [9, 13]. Furthermore, if more than 100 g of skeletal muscle is damaged, the serum haptoglobin binding capacity becomes saturated with myoglobin and remaining free circulating myoglobin can precipitate and obstruct the renal tubules, leading to tubular necrosis, and development of acute kidney injury (AKI).

About 7-10% of all cases of AKI in United States are contributable to RB [14]. Chatzizisis et al explored the RB complications that included life-threatening disorders such as hyperkalemic cardiotoxicity, acute renal failure, and early hy-
pocalcemia [15]. Irrespective of the cause of RB, the mortality rate may still be as high as 8% with an RB-induced AKI mortality rate of approximately 20% [16].

Case Report

A 20-year-old young male with no significant past medical history presented to the emergency room with severe fatigue and pain all over his body with reddish brown urine following 2 h of intense workout at the gym. He was well hydrated during his workout session. He denied any history of trauma, chest pain, breathlessness or use of illicit drugs.

He lives an active lifestyle and engages in school sports like football and basketball on regular basis. He occasionally takes aspirin for body pain and aches. Two days prior to admission, he spent 2 h at the gym specifically doing upper body workout, carrying 35 - 75 lbs. He has not been to the gym for the last 5 years. On the first day, he tolerated the workout well; however, on the second day after working out for an hour in the gym, he immediately felt exhausted and fatigued, but forced himself to continue for another hour. He was not on any workout supplements or weight loss products. After his workout session, he experienced intense diffuse muscle pain, which restricted his movement. He took two tablets of Aleve that mildly alleviated the pain. Four hours later, he started to pass reddish brown urine and was brought to the emergency department.

Vitals on presentation revealed normal heart rate of 118 beats per minute, blood pressure of 126/78 mm Hg, and respiratory rate of 20 breaths per minute and he was afebrile. On examination, he was well oriented to time, place and person. Lung and heart examinations were normal. Abdominal was soft and non-tender. Musculoskeletal system examination revealed muscle pain with a severity of 8/10 and tenderness in arms, back and shoulders with intact reflexes.

His initial set of ER lab showed an elevated hemoglobin of 16.7 g/dL with a white cell count of 13,440/mm³. The CK level on admission was about 510,924 IU/L. Creatinine (Cr) was 1.33 mg/dL with blood urea nitrogen (BUN) of 20 mg/dL, AST was 1,951 IU/L, ALT was 381 IU/L, anion gap was 11.0 mEq/L, albumin was 4.5 g/dL, and inorganic phosphate was 5 mg/dL. His urine analysis showed specific gravity > 1.030 and protein > 300 mg/dL, blood and bilirubin was detected as high, and nitrite was positive with moderate leukocytes. RBCs casts were present with mucous threads and the urine myoglobin level > 12,700 μg/L (reference range: < 28 μg/L). Drug screen turned out to be negative.

An initial diagnosis of RB was made and the patient was bolused on 3 L of sodium chloride 0.9%. Muscle aches were managed by I/V morphine 2 mg/1 mL PRN. Chest X-ray, EKG and ECHO were normal with no evidence of any cardiopulmonary disease. His renal ultrasound showed no hydronephrosis, with increased echogenicity of renal cortices, suggestive of medical renal disease with a trace of lateral perinephric fluid.

He was admitted to the ward for further management. Urine output and renal plasma flow were closely monitored. Fluids were continued, titrating to 250 mL/h. The CK was repeated and the count dropped to 200,000 IU/L. Overnight, the patient developed stage 3 AKI as his creatinine level started to rise from 1.33 mg/dL on admission to 7 mg/dL, but his electrolytes were normal with no acid base disorder and adequate urine output.

Patient was put on renal diet with low potassium and low phosphorus with protein restriction to about 60 - 75 g/day (0.8 - 1 g/kg IBW according to AKI guidelines) and also total calorie intake of about 2,250 kcal/kg (30 kcal/kg IBW according to
AKI guidelines). However, the S. Cr continued to rise, peaking to 13.88 mg/dL on day 10. Further blood work revealed hyperphosphatemia reaching 9.5 mg/dL. As expected, calcium levels started to trend downwards from 9.4 to 6.9 mmol/L. Phosphorus binder, Renagel was started. Given the young age of the patient and the permissible levels of potassium with adequate urine output, dialysis was held off and the patient was diligently monitored and treated empirically for an AKI secondary to RB.

The CK and inorganic phosphate trended downwards steadily. It is worth noting that the patient had fluctuating BP acutely. Efforts were made to rule out other etiologies. ANA, ANTI DNA Ds, HIV, and thyroid panel revealed negative results (Fig. 1).

The patient was counseled and discharged on request. He was discharged on famotidine 20 mg tab, and sevelamer HCL 800 mg and sennosides 8.6 mg tab. On discharge, his CK level was 1,701 IU/L, and phosphorus was 6.9 mg/dL with a Cr of 12.64 mg/dL and a BUN of 91 mg/dL. His follow-up appointment with the primary and nephrologists reveal a CK of 338 IU/L and a Cr of 1.99 mg/dL and the patient made a complete recovery.

Discussion

The median age of RB is 47 years; however, the incidence and risk factors for RB in younger patients have not been established [17]. A large number of risk factors are known to cause RB and can be categorized as intrinsic muscle dysfunction, use of prescribed medications such as amphetamines and phen-cyclidine hydrochloride, use of illicit drugs such as heroin, metabolic disorders, trauma, heatstroke, infections, seizure or long-term immobilization [1]. In a retrospective study of RB in young patients, Chen et al reported 87% were males with trauma (26.7%) and infections (18.3%) were the most common causes [18].

The association of excessive physical exertion (e.g. marathoners and military recruits) with RB is well documented in the medical literature [19]. Occurrence of RB depends on the overall fitness of the body including the targeted muscles and the intensity of physical activity performed. In 2016, Hummel et al presented a series of cases of RB in young athletes. They described numerous cases of RB caused by one or a few strenuous workout sessions after a period of prolonged inactivity [20]. This article described six cases in which patients presented with RB after participation in the CrossFit exercise regimen. These cases are significant as three of the cases involved patients that had been participating in CrossFit for months to years. This disputes the assumption that exertional RB solely occurs when exerting oneself after a prolonged period of immobility.

Often multiple confounding risk factors are present. Many cases involved the athletes admitting to poor fluid intake prior to exercising. Many athletes admitted to consuming caffeine, protein substances, and NSAIDs that may contribute to the risk of RB. Other cases involved patients suffering from heat stroke during the exercise session. In 2012, Oh et al reported on a cluster of cases of acute exertional RB precipitated by football camp resistance exercises performed in a hot building. The athletes were given little time to re hydrate, which may have led to further worsening of their condition. The authors attribute poor knowledge about risk factors and preventative strategies regarding RB [21]. Knowledge of the true prevalence of exercise-induced RB can help better educate athletic training programs. In the younger demographic population, in addition to exertional RB, it is important to consider infection as a possible cause of RB.

Some level of CK rise is normal in exercising individuals. O’Brien et al studied serum CK levels in military recruits in training. They found that patients with exertional RB had a higher average S. CK level (40,471 IU/L) than other etiologies and had a lower incidence of AKI. They found these patients responded very well to rehydration. Despite peak serum CK levels >100,000 IU/L in some cases, acute renal failure did not often occur, and this form of RB seemed very responsive to I/V hydration.

Conclusion

We present the case of severe RB in a young adult that occurred just after a 2-hour workout session at the gym. His initial CPK at time of presentation was unusually high reaching over 500,000 IU/L. Despite getting aggressively bolused with IV fluids, there was rapid deterioration in kidney function and the patient advanced into AKI reaching a peak Cr of 13.88 mg/dL. All measures were taken to closely monitoring of renal functions and CK levels. As the patient was stable with adequate urine output and no acid base disorder, we decided to treat him with IV fluids only and renal replacement therapy held off. By day 10, the S. Cr started to plateau and then decline. Our patient has made a complete recovery and to date is healthy.

Our patient’s course of gradual recovery from stage 3 AKI secondary to severe Rb with abnormally elevated CK highlights the paramount importance of early IV fluids intervention and clinicians should also bear in mind the patient’s overall health status before considering renal replacement therapy. More research especially in the younger population is needed to better guide clinicians in their treatment protocols.

Consent

Written informed consent was obtained from the patient for publication of this case report.

References