REVIEW ARTICLE

CURRENT CONCEPTS

Rhabdomyolysis and Acute Kidney Injury

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HABDOMYOLYSIS — LITERALLY, THEDISSOLUTION OF STRIPED (SKELETAL) muscle — is characterized by the leakage of muscle-cell contents, including electrolytes, myoglobin, and other sarcoplasmic proteins (e.g., creatine kinase, aldolase, lactate dehydrogenase, alanine aminotransferase, and aspartate amino transferase) into the circulation. Massive necrosis, which is manifested as limb weak ness, myalgia, swelling, and, commonly, gross pigmenturia without hematuria, is the common denominator of both traumatic and nontraumatic rhabdomyolysis. Acute kidney injury is a potential complication of severe rhabdomyolysis, regardless of whether the rhabdomyolysis is the result of trauma or some other cause, and the prognosis is substantially worse if renal failure develops. In contrast, in less severe forms of rhabdomyolysis or in cases of chronic or intermittent muscle destruction — a condition sometimes called hyperCKemia — patients usually present with few symptoms and no renal failure. We review the pathophysiological characteristics and management of acute kidney injury associated with rhabdomyolysis.

There are eight commonly reported categories of rhabdomyolysisTable 1). Exogenous agents that can be toxic to muscles, especially alcohol, illicit drugs, and lipid-lowering agents, are common nontraumatic causes. Recurrent episodes of rhabdo myolysis are often a sign of an underlying defect in muscle metabolism^{1,3,4}

Acute rhabdomyolysis occasionally develops in patients with structural myopa thies when they are performing strenuous exercise, are under anesthesia, have taken drugs that are toxic to muscles, or have viral infections.¹ When a diagnosis of acute rhabdomyolysis is suspected, histochemical, immunohistochemical, and mito chondrial respiration studies performed on a muscle-biopsy specimen may yield a specific diagnosis. It is important to wait several weeks or months after the clinical event to perform a biopsy, because the results of a biopsy will typically be uninfor mative at an early stage. Thus, the specimen may appear normal or show no spe cific findings other than necrosis during and early after the acute episode of rhabdomyolysis (Fig. 1)?.5

The mechanisms involved in the pathogenesis of rhabdomyolysis are direct sarcolemmic iniury (e.g., trauma) or depletion of ATP within the myocyte, leading to an unregulated increase in intracellular calcium. Sarcoplasmic calcium is strictly regulated by a series of pumps, channels, and exchangers that maintain low levels of calcium when the muscle is at rest and allow the increase that is necessary for actin—myosin binding and muscle contraction. Depletion of ATP impairs the function of these pumps, resulting in a persistent increase in sarcoplasmic calcium that leads to persistent contraction and energy depletion and the activation of calcium-dependent neutral proteases and phospholipases; the result is the eventual destruction of myofibrillar, cytoskeletal, and membrane proteins, followed by lysosomal digestion of fiber contents. Ultimately, the myofibrillar network breaks down, resulting in disintegration of the myocyte? In the case of patients with rhabdomy

Table 1. Major Categories and Commonly Reported Causes of Rhabdomyolysis. **Commonly Reported Cause** Category Trauma Crush syndrome Exertion Strenuous exercise, seizures, alcohol withdrawal syndrome Muscle hypoxia Limb compression by head or torso during prolonged immobilization or loss of consciousness,* major artery occlusion Genetic defects Disorders of glycolysis or glycogenolysis, including myophosphorylase (glycogenosis type V), phosphofructokinase (glycogenosis type VII), phosphorylase kinase (glycogenosis type VIII), phosphoglycerate kinase (glycogenosis type IX), phosphoglycerate mutase (glycogenosis type X), lactate dehydrogenase (glycogenosis type XI) Disorders of lipid metabolism, including carnitine palmitoyl transferase II, longhain acyl CoA dehydrogenase, short-chain L-3-hydroxyacyl-CoA dehydrogenase, medium-chain acyl-CoA dehydrogenase, very-long-chain acyl-CoA dehydrogenase, medium-chain 3-ketoacyl-CoA, thiolase† Mitochondrial disorders, including succinate dehydrogenase, cytochrome c oxidase, coenzyme Q10 Pentose phosphate pathway: glucose-6-phosphate dehydrogenase Purine nucleotide cycle: myoadenylate deaminase Infections:: Influenza A and B, coxsackievirus, Epstein-Barr virus, primary human immunodeficiency virus, legionella Streptococcus pyogenes, Staphylococcus aureus (pyomyositis), clostridium Body-temperature changes Heat stroke, malignant hyperthermia, malignant neuroleptic syndrome, hypothermia Metabolic and electrolyte disorders Hypokalemia, hypophosphatemia, hypocalcemia, nonketotic hyperosmotic conditions, diabetic ketoacidosis Drugs and toxins Lipid-lowering drugs (fibrates, statins), alcohol, heroin, cocaine

Idiopathic (sometimes recurrent)

olysis caused by trauma, additional injury results persons who had undergone trauma than among from ischemia reperfusion and inflammation by persons with muscle disease, and the incidence neutrophils that infiltrate damaged muscle?

EPIDEMIOLOGY OF MYOGLOBINURIA-INDUCED ACUTE KIDNEY INJURY

Acute kidney injury associated with myoglobinu tions and clinical scenarios. The reported incidence ranges from 13% to approximately 50%.-11 of acute kidney injury was 46%. Although rhabout present. Although rhabout present. Although rhabout present. Although rhabout present. used illicit drugs or abused alcohol and among acute kidney injury recover renal function!4

was particularly high among persons with more than one recognized causal factor.10

The outcome of rhabdomyolvsis is usually good provided that there is no renal failure. Nevertheless, mortality data vary widely according to the study population and setting and the ria is the most serious complication of both trau number and severity of coexisting conditions. In matic and nontraumatic rhabdomyolysis, and it a study in which the incidence of vasculopathy may be life-threatening. Acute kidney injury as a leading to rhabdomyolysis as a result of limb is complication of rhabdomyolysis is quite common, chemia was high, the overall mortality was 32%? representing about 7 to 10% of all cases of acute In contrast, the study by Melli et al. of hospital kidney injury in the United States^{4,9} The true in- ized patients, in whom the abuse of illicit drugs cidence of acute kidney injury in rhabdomyolysis and alcohol was the most frequently identified is difficult to establish owing to varying definicause of rhabdomyolysis, showed a mortality of 3.4% among patients with acute kidney injury.10 Among patients in the intensive care unit, the In a study by Melli et al. involving 475 hospital mortality has been reported to be 59% when ized patients with rhabdomyolysis, the incidence acute kidney injury is present and 22% when it is domyolysis from any cause can lead to acute kid tients with rhabdomyolysis and acute kidney in ney injury, in this study, the incidence of acute jury is reported to be close to 80%, and the ma kidney injury was higher among persons who jority of patients with rhabdomyolysis-induced

^{*} Rhabdomyolysis from this cause is associated with a crush syndrome-like mechanism.

[†] CoA denotes coenzyme A.

[‡] In most cases, the mechanism is unclear.

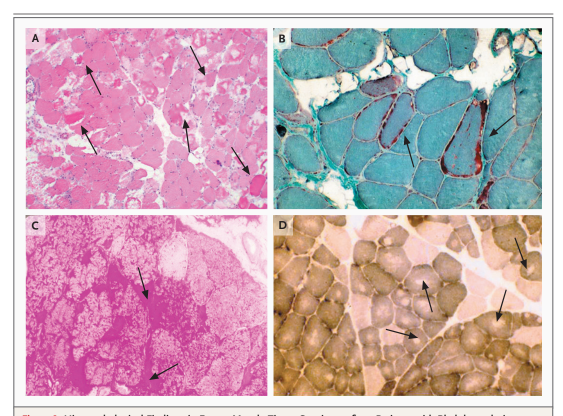


Figure 1. Histopathological Findings in Frozen Muscle-Tissue Specimens from Patients with Rhabdomyolysis. Panel A shows massive muscle necrosis (arrows) in a patient with statinrelated rhabdomyolysis (hematoxylin and eosin). This histologic feature would be similar in every case of rhabdomyolysis, irrespective of the cause. Panel B shows the typical ragged-red fibers (arrows) in a musclebiopsy specimen from a patient with mitochondrial myopathy that was obtained 3 months after an episode of severe rhabdomyolysis. The mitochondrial dysfunction was confirmed by a mitochondrial respiratory chain-based assay (Gomori's trichrome). Panel C shows periodic acid-Schiff (PAS)positive material (arrows) in some muscle fibers in a case of McArdle's disease. The biopsy was performed a few months after the patient's recovery from recurrent rhabdomyolysis (PAS stain). Panel D shows a muscle-biopsy specimen from a patient with central core disease. The specimen was obtained after the patient's recovery from malignant hyperthermia. Abundant central cores can be seen (arrows) (NADH-tetrazolium reductase stain).

PATHOGENESIS OF MYOGLOBIN-INDUCED ACUTE KIDNEY INJURY

Myoglobinuria occurs only in the context of rhab domyolysis. Myoglobin is a dark red 17.8-kDa pro constriction, and it precipitates when it interacts tein that is freely filtered by the glomerulus, enters with the Tamm-Horsfall protein, a process fathe tubule epithelial cell through endocytosis, and vored by acidic urine. Tubule obstruction occurs is metabolized. It appears in the urine only when principally at the level of the distal tubules, and the renal threshold of 0.5 to 1.5 mg of myoglobin direct tubule cytotoxicity occurs mainly in the per deciliter is exceeded and is grossly visible as proximal tubules. reddish-brown ("tea-colored") urine when serum myoglobin levels reach 100 mg per decilite^{‡5}; toxic effect in the tubules unless the urine is therefore, not all cases of rhabdomyolysis are as acidic. Myoglobin is a heme protein; it contains sociated with myoglobinuria.

rhabdomyolysis impairs the glomerular filtration molecular oxygen can promote the oxidation of rate are unclear, experimental evidence suggests Fe2+ to ferric oxide (Fe3+), thus generating a hv

chemic tubule injury, and tubular obstruction all play a role (Fig. 2).16 Myoglobin becomes concentrated along the renal tubules, a process that is enhanced by volume depletion and renal vaso

Myoglobin seems to have no marked nephro iron, as ferrous oxide (Fe²⁺), which is necessary Although the exact mechanisms by which for the binding of molecular oxygen. However, that intrarenal vasoconstriction, direct and is- droxyl radical. This oxidative potential is counter acted by effective intracellular antioxidant mole as 5000 U per liter, this usually occurs when co leads to uncontrolled leakage of reactive oxygen and acidosis are present¹¹ For example, in patients species, and free radicals cause cellular injury. It with chronic myopathies such as muscular dys has been suggested that heme and free iron- trophies and inflammatory myopathies, acute kid driven hydroxyl radicals are critical mediators of ney injury seldom develops unless a superim deferoxamine (an iron chelator) and glutathiones myopathies, on the other hand, may have moder itself can exhibit peroxidase-like enzyme activity but not overt myoglobinuria²⁴ Myoglobinuria can that leads to uncontrolled oxidation of biomole be inferred if urinary dipstick testing shows a isoprostanes.19

Renal vasoconstriction is a characteristic feature of rhabdomyolysis-induced acute kidney in jury and is the result of various combinations of several mechanisms. First, intravascular volume depletion due to fluid sequestration within dam aged muscle promotes homeostatic activation of (Table 2).²⁵ Myoglobin is the true pathogenic fae the renin-angiotensin system, vasopressin, and the sympathetic nervous system. Second, experi mental studies have shown that there are additional vascular mediators in the reduction of renal blood flow, including endothelin-1, throm boxane A_2 , tumor necrosis factor α ; and F_2 -isoprostanes^{9,20}; a deficit in the vasodilator nitric oxide, which can be attributed to the scavenging spleen).²⁶ Therefore, measurement of serum myo has also been shown to be a mediator in the re duction in renal blood flow.16 Collectively, these vascular mediators appear to be locally stimulated by oxidant injury and leukocyte-mediated inflammation as a result of the endothelial dys kidney injury.21

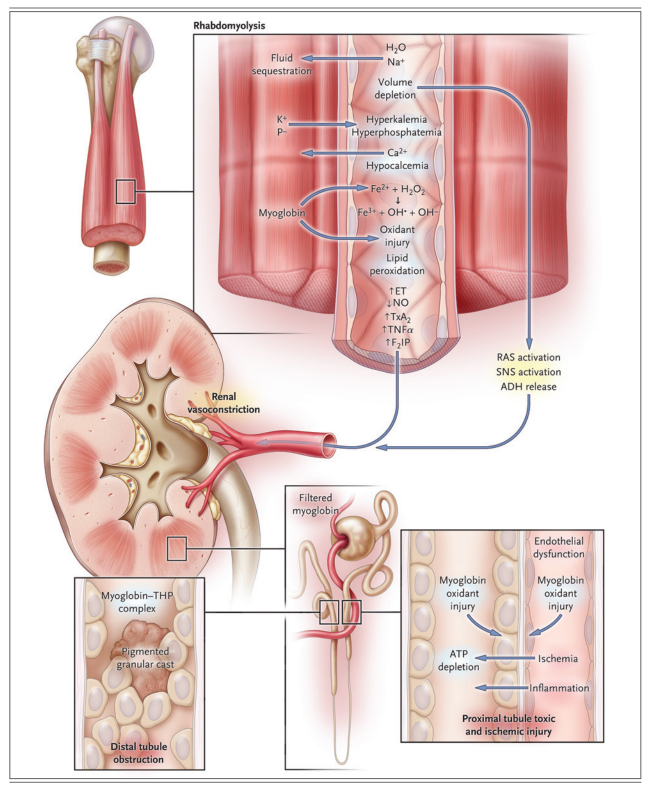
RENAL MANIFESTATIONS OF RHABDOMYOLYSIS

Patients with acute rhabdomyolysis usually pixent with pigmented granular casts, reddish-brown urine supernatant, and markedly raised serum creatine kinase. There is no defined threshold sis-induced acute kidney injury that is different value of serum creatine kinase above which the from the manifestation of other forms of acute risk of acute kidney injury is markedly increased, tubular necrosis is the frequent, but not universal. A very weak correlation between the peak creatine presence of a low fractional excretion of sodium kinase value and the incidence of acute kidney (≤1%), perhaps reflecting the primacy of pre injury or peak serum creatinine has been report glomerular vasoconstriction and tubular occlued. 10,11,22 The risk of acute kidney injury in rhab sion rather than tubular necrosis? The fractional domyolysis is usually low when creatine kinase excretion of sodium is a measurement of the levels at admission are less than 15,000 to 20,000 U percentage of filtered sodium that is excreted in per liter.^{12,13,23} Although acute kidney injury may the urine, and low levels in patients with acute

cules. However, cellular release of myoglobin existing conditions such as sepsis, dehydration, tubule damage owing to the protective effects of posed event is present. Patients with these chronic More recently, it has been shown that myoglobin ately raised concentrations of plasma myoglobin cules, lipid peroxidation, and the generation of positive result for blood when there are no red cells in the sediment. This false positive result for blood occurs because the dipstick test is un able to distinguish between myoglobin and hemo globin. The test has a sensitivity of 80% for the detection of rhabdomyolvsis. Other causes of pig mented urine should be taken into consideration tor in rhabdomyolysis-induced acute kidney injury but is seldom measured directly in urine or plasma. Serum myoglobin levels peak well before serum creatine kinase levels, and serum myoglobin has a rapid and unpredictable metabolism, which func tions partly through the kidney but mainly out side the kidney (probably through the liver or effect of myoglobin in the renal microcirculation, globin has a low sensitivity for the diagnosis of rhabdomyolysis.27

Acute kidney injury associated with rhabdo myolysis often leads to a more rapid increase in plasma creatinine than do other forms of acute kidney injury. However, this finding may reflect function that is common to other forms of acute the overrepresentation of young, muscular men among patients with rhabdomyolysis rather than increased creatinine or creatine release from in jured muscle:14,28,29 Similarly, a low ratio of blood urea nitrogen to creatinine is often seen in patients with rhabdomyolysis. Rhabdomyolysis-induced acute kidney injury frequently causes oliguria and occasionally causes anuria.

Another characteristic feature of rhabdomyoly be associated with creatine kinase values as low kidney injury are an indication of the relative



integrity of tubular functions. However, when is chemic or toxic acute tubular necrosis is established, both urinary sodium and the fractional excretion of sodium are raised.

Electrolyte abnormalities that occur as a result of the release of cellular components often accompany and determine the severity of rhab domyolysis-induced acute kidney injury. Because

Table 2 Causes and Microscopic Features of Red and Brown Urine

Table 2. Causes and Microscopic Features of Red and Brown Offine.					
Cause	Results of Test for Blood in Fresh Urine*	Sediment†‡	<u>Supernatant÷</u>		
<u>Hematuria</u>	+ to ++++	Red	Yellow		
<u>Myoglobinuria</u>	+ to ++++	Normal	Red to brown		
<u>Hemoglobinuria</u>	+ to ++++	Normal	Red to brown		
Porphyria	Negative	Normal	Red		
Bile pigments	Negative	Normal	Brown		
Food and drugs§	Negative	Normal	Red to brown		

^{*} Urine was tested with the use of a dipstick test. This is a semiquantitative test of the number of erythrocytes per microliter. Results range from + (5 to 10 erythrocytes per microliter) to ++++ (approximately 250 erythrocytes per microliter). † Normal refers to white or yellow in color, unremarkable in the absence of cells, crystals, or cylinders.

they may precede the acute kidney injury, elee trolyte levels should be measured as soon as rhab domyolysis is diagnosed. The electrolyte abnor malities that can occur with rhabdomyolysis include hyperkalemia (which can be rapidly in creasing), hyperphosphatemia, hyperuricemia, high anion-gap metabolic acidosis, and hypermagnesemia mainly when renal failure is present.4,15,22,31 High levels of phosphate can bind to urine. calcium, and deposition of calcium-phosphate hyperphosphatemia inhibits 1α -hydroxylase, thus limiting the formation of calcitriol (1,25-dihy droxyvitamin D₃), the active form of vitamin D. Hyperkalemia is an early manifestation of rhab

Figure 2 (facing page). Pathophysiological Mechanisms in Rhabdomyolysis-Induced Acute Kidney Injury.

Fluid sequestration in injured muscle induces volume depletion and consequent activation of the sympathetic nervous system (SNS), antidiuretic hormone (ADH), and the renin-angiotensin system (RAS), all of which favor vasoconstriction and renal salt and water conservation. In addition, myoglobininduced oxidative injury increases vasoconstrictors and decreases vasodilators. Kidney injury results from a combination of ischemia due to renal vasoconstriction, direct tubular toxicity mediated by myoglobin-associated oxidative injury (inset, lower right), tubular damage due to ischemia, and distal tubule obstruction due to precipitation of the Tamm-Horsfall protein-myoglobin complex (inset, lower left) in addition to sloughed tubular cells forming cellular cast. As in acute kidney injury due to other causes, en dothelial dysfunction and local inflammation contribute to tissue damage and organ dysfunction. ET denotes endothelin, F2 IP F2 isoprostanes, NO nitric oxide, THP Tamm–Horsfall protein, TNF- α tumor necrosis factor α , TxA_2 thromboxane A_2 , and VC vasoconstriction.

domyolysis, and serum potassium can occasion ally reach <u>life-threatening</u> levels both in patients with severe traumatic rhabdomyolysis and in those with nontraumatic rhabdomyolysis:15,31 Hyperuricemia is also usually present owing to the liberation of nucleosides from injured muscle and can contribute to renal tubule obstruction since uric acid is insoluble and may precipitate in acidic

Hypocalcemia is a common complication of complexes in soft tissues can occur. In addition, rhabdomyolysis and usually results from calcium entering the ischemic and damaged muscle cells and from the precipitation of calcium phosphate with calcification in necrotic muscle. Hypercal cemia associated with recovery of renal function is unique to rhabdomyolysis-induced acute kid ney injury and results from the mobilization of calcium that was previously deposited in muscle, the normalization of hyperphosphatemia, and an increase in calcitriol.32

TREATMENT AND PREVENTION

Patients with rhabdomyolysis that is associated with acute kidney injury usually present with a clinical picture of volume depletion that is due to the sequestration of water in injured muscles. Therefore, the main step in managing the condi tion (Table 3) remains the early, aggressive reple tion of fluids; patients often require about 10 liters of fluid per day,31 with the amount administered depending on the severity of the rhabdo myolysis.33 There are no randomized trials that have evaluated fluid repletion in patients with the crush syndrome resulting from injuries sus

The sediment and supernatant were examined after centrifugation of 10 to 15 ml of urine at 1500 to 3000 rpm for 5 minutes.

Food and drugs that can cause red urine include beets, blackberries, rhubarb, food coloring, fava beans, phenolphthalein, rifampin, doxorubicin, deferoxamine, chloroquine, ibuprofen, and methyldopa. Those that can cause brown urine include levodopa, metronidazole, nitrofurantoin, iron sorbitol, chloroquine, and methyldopa.

Table 3. Steps in the Prevention and Treatment of Rhabdomyolysis-Induced Acute Kidney Injury.

Check for extracellular volume status, central venous pressure, and urine output.*

Measure serum creatine kinase levels. Measurement of other muscle enzymes (myoglobin, aldolase, lactate dehydrogenase, alanine aminotransferase, and aspartate aminotransferase) adds little information relevant to the diagnosis or management.

Measure levels of plasma and urine creatinine, potassium and sodium, blood urea nitrogen, total and ionized calcium, magnesium, phosphorus, and uric acid and albumin; evaluate acid-base status, blood-cell count, and coagulation.

Perform a urine dipstick test and examine the urine sediment.

Initiate volume repletion with normal saline promptly at a rate of approximately 400 ml per hour (200 to 1000 ml per hour depending on the setting and severity), with monitoring of the clinical course or of central venous pressure.

Target urine output of approximately 3 ml per kilogram of body weight per hour (200 ml per hour).

Check serum potassium level frequently.

Correct hypocalcemia only if symptomatic (e.g., tetany or seizures) or if severe hyperkalemia occurs.

Investigate the cause of rhabdomyolysis.

Check urine pH. If it is less than 6.5, alternate each liter of normal saline with 1 liter of 5% dextrose or 0.45% saline. plus 100 mmol of bicarbonate. Avoid potassium and lactate-containing solutions.

Consider treatment with mannitol (up to 200 g per day and cumulative dose up to 800 g). Check for plasma osmolality and plasma osmolal gap. Discontinue if diuresis (>20 ml per hour) is not established.

Maintain volume repletion until myoglobinuria is cleared (as evidenced by clear urine or a urine dipstick testing result that is negative for blood).

Consider renal-replacement therapy if there is resistant hyperkalemia of more than 6.5 mmol per liter that is symptomatic (as assessed by electrocardiography), rapidly rising serum potassium, oliguria (<0.5 ml of urine per kilogram per hour for 12 hours), anuria, volume overload, or resistant metabolic acidosis (pH < 7.1).

patients in whom acute kidney injury did not de tial hypocalcemic phase of rhabdomyolysis. velop (Table 4).8,23,31,33,39 Therefore, early, aggres the crush syndrome.34,35

Although the need for volume repletion is es tablished, the composition of the fluid used for repletion remains controversial. Some investiga tors recommend administering sodium bicarbon ate, which results in an alkaline urine, as first proposed by Bywaters and Beall, 8,39,40 whereas others argue against this approach and favor nor mal or 0.45% saline solution. The three empirical advantages of alkalinization that have been rhabdomyolysis. First, it is known that precipita complex is increased in acidic urine.¹⁷ Second, alkalinization inhibits reduction-oxidation (re in rhabdomyolysis, thus ameliorating tubule in for dialysis, or death in the sample as a whole, jury.41 Third, it has been shown that metmyo although the results suggested that it might be

tained in a disaster such as an earthquake. How medium in the isolated perfused kidney. The ever, most, if not all, reports show that patients principal, and probably the only, disadvantage of in whom acute kidney injury developed had a lon alkalinization is the reduction in ionized calcium, ger delay in receiving supportive therapy than did which can exacerbate the symptoms of the ini-

The clinical benefits of alkalinization as com sive volume repletion is crucial in patients with pared with simple volume repletion are not firm ly established. Comparative studies usually have small sample sizes and show a combination of measures (e.g., alkalinization plus mannitol) that preclude an analysis of the effectiveness of the particular single measure34-38 (Table 4). In one study, renal outcomes did not differ significant ly between patients treated with bicarbonate plus mannitol and those treated with saline alone, although peak serum creatine kinase values were below 5000 U per liter, a finding indicating that noted are based on studies in animal models of the degree of injury was mild, making treatment effect difficult to appreciate³⁶ In the largest study tion of the Tamm-Horsfall protein-myoglobin of patients who had undergone trauma (2083 patients), rhabdomyolysis developed in 85% of the patients, and administration of bicarbonate plus dox) cycling of myoglobin and lipid peroxidation mannitol did not prevent renal failure, the need globin induces vasoconstriction only in an acidic beneficial in patients with peak creatine kinase

^{*} In the case of the crush syndrome (e.g., earthquake, building collapse), institute aggressive volume repletion promptly before evacuating the patient.

Study	Study Design	Patient Group	No. in Sample	Therapeutic Strategy	Outcome in Patients with Acute Kidney Injury
Shimazu et al. ³⁴	Retrospective	Patients with the crush syndrome	14	Late vs. early initiation of therapy; high (>10 liters for 48 hours) vs. low volume of hydration	Better if therapy initiated early high volume of hydration better
Gunal et al. ³⁵	Retrospective	Patients with the crush syndrome	16	Early vs. late treatment with nor- mal saline followed immedi- ately by bicarbonate	Better if treatment initiated early
Homsi et al. ³⁶	Retrospective	Patients in the intensive care unit	24	Normal saline vs. normal saline plus bicarbonate and mannitol	No difference
Brown et al. ³⁷	Retrospective	Patients with trauma	2083	Normal saline vs. bicarbonate plus mannitol	No difference
Cho et al. ³⁸	Prospective, randomized	Patients with intoxication from doxylamine	28	Ringer's lactate vs. normal saline; bicarbonate if urine pH is <6.5	No effect on peak creatine ki- nase level or recovery with Ringer's lactate as com- pared with normal saline; more bicarbonate needed with normal saline than with Ringer's lactate

values of more than 30.000 U per liter.³⁷ In a uted to doxylamine intoxication, 28 patients were ic diuretic, it increases urinary flow and the atine kinase levels were less than 10,000 U per it is a free-radical scavenger^{4,8,20} Most data on liter, and it appears that acute kidney injury did the action of mannitol come from studies in ani data were not reported. Whatever the real, consis effect of mannitol may be attributable to its os sive infusion of normal saline alone can contrib has supported the evidence-based use of man ute to metabolic acidosis, mainly owing to the nitol, and some clinical studies suggest no bene dilution of serum bicarbonate with a solution ficial effects.^{36,37} In addition, high accumulated relatively high in chloride ions, generating hyper doses of mannitol (>200 g per day or accumu Therefore, administration of both normal saline and tubular toxicity, a condition known as osand sodium bicarbonate seems to be a reason motic nephrosis. 45,46 However, many experts con with metabolic acidosis (Table 3). If sodium bi kidney injury and relieve compartmental presnormal saline.

The use of diuretics remains controversial, but randomized, prospective trial of fluid repletion it is clear that it should be restricted to patients with Ringer's lactate as compared with normal in whom the fluid repletion has been achieved. saline in patients with rhabdomyolysis attrib- Mannitol may have several benefits; as an osmot randomly assigned to receive one of the solu flushing of nephrotoxic agents through the renal tions.38 Sodium bicarbonate was added in both tubules; as an osmotic agent, it creates a gradient groups if the urine pH was less than 6.5 after 12 that extracts fluid that has accumulated in injured hours of aggressive volume repletion. Peak cre muscles and thus improves hypovolemia; finally, not develop in any of the patients, although these mals, which collectively show that the protective tent benefits of urine alkalinization in patients motic diuretic action rather than to the other with rhabdomyolysis, there is evidence that mas mechanisms.⁴⁴ No randomized, controlled trial chloremic metabolic acidosis with observed re lated doses of >800 g) have been associated with ductions in serum pH of as much as 0.30 units^{4,3} acute kidney injury due to renal vasoconstriction able approach when fluid is being replenished in tinue to suggest that mannitol should be used to patients with rhabdomyolysis, especially patients prevent and treat rhabdomyolysis-induced acute carbonate is used, urine pH and serum bicar sure.^{20,45-47} During the time mannitol is being bonate, calcium, and potassium levels should be administered, plasma osmolality and the osmolal monitored, and if the urine pH does not rise gap (i.e., the difference between the measured after 4 to 6 hours of treatment or if symptom and calculated serum osmolality) should be mon atic hypocalcemia develops, alkalinization should itored frequently and therapy discontinued if ade be discontinued and hydration continued with quate diuresis is not achieved or if the osmolal gap rises above 55 mOsm per kilogram⁴⁶ Loop

Table 5. Approach to the Management of Hyperkalemia (Serum Potassium ≥5.5 mmol per Liter) in Rhabdomyolysis.

Check for potassium levels every 4 hours in cases of severe rhabdomyolysis (creatine kinase level >60,000 to 80,000 U per liter) or suspected systemic toxin. Treat rapidly rising potassium levels aggressively.

Obtain an electrocardiogram and check for severe manifestations (QRS interval widening, small P waves, severearrhythmias thought to be caused by high levels of potassium). Consider cardiac monitoring and admission to intensive care unit if the potassium level is higher than 6 mmol per liter, if there are abnormalities on the electrocardiogram, or if rhabdomyolysis is severe, with rapidly rising potassium.

Check for plasma calcium levels. Hypocalcemia seriously aggravates the adverse electrical effects of hyperkalemia.

If the electrocardiogram shows severe irregularities, administer calcium chloride or calcium gluconate by intravenous infusion. Consider slow continuous infusion if hypocalcemia is present. Anticipate possible hypercalcemia in late rhabdomyolysis. Do not mix with bicarbonate solutions.

If potassium level is higher than 6 mmol per liter, shift potassium into cells. Serum potassium will be lowered approximately 10 to 30 minutes after the following measures are performed, and the effect will last for 2 to 6 hours.

Administer insulin and glucose by means of a slow intravenous push; monitor glucose with the use of fingerstick testing.

Administer a β_2 -adrenergic agonist such as albuterol, 10 to 20 mg in 4 ml of normal saline by inhalation of aerosol over 10 minutes. Do not use as a single measure; combine with glucose and insulin for additive effect.

Administer sodium bicarbonate if the patient has acidemia. This treatment may worsen the manifestations of hypcalcemia, and the efficacy is not as consistent as that with insulin and glucose or albuterol. Do not use as a single measure.

Remove potassium from the body with the use of either resins or dialysis as indicated; the use of diuretics is optional.

Administer cation-exchange resin (sodium polystyrene sulfonate) orally or as a retention enema (avoid sorbitol in such cases and avoid after surgery).

Perform hemodialysis if the above measures fail or if severe renal failure or severe hyperkalemia develops. Consider hemodialysis when rhabdomyolysis is associated with marked tissue breakdown and rapidly rising serum potassium levels. Check serum potassium levels 4 hours after hemodialysis, since a rebound increase can occur. Previous measures of potassium shift into cells may decrease the efficiency of hemodialysis with respect to emoval of potassium.

Administer loop diuretics such as furosemide, but only after the patient's fluid level has been expanded.

crease the risk of myoglobin precipitation, but no cated, principally with intermittent hemodialy study has shown a clear benefit in patients with sis, which can correct electrolyte abnormalities rhabdomyolysis. Therefore, loop diuretics in rhab rapidly and efficiently. 8,9,47 Conventional hemodomyolysis-induced acute kidney injury should be dialvsis does not remove mvoglobin effectively used in the same manner as that recommended in owing to the size of the protein and is therefore

be treated promptly; the correction of hyper- ventive extracorporeal elimination has been stud the disease, is especially important (Table 5).49 to have no effect on outcomes or on the myo-Agents that cause a shift of potassium from the globin burden of the kidneys; o continuous venotonic glucose and bicarbonate) are effective only shown some efficacy in removing myoglobin, potassium from the body is diuresis (effective ka and high volumes of ultrafiltration (convection). or dialysis, 4,8,9,15 In contrast, early hypocalcemia case reports, and the effect on outcomes is un containing chelators should be used with caution differ significantly between patients who are could increase the precipitation of calcium phos tinuous venovenous hemodiafiltration.²⁷ Until phate in injured muscle.^{4,8,9,15}

When acute kidney injury is severe enough to hemofiltration cannot be recommended. produce refractory hyperkalemia, acidosis, or vol

diuretics also increase urinary flow and may de ume overload, renal-replacement therapy is indi acute kidney injury that is due to other causes. 47,48 usually mandated by renal indications. However, The electrolyte abnormalities associated with owing to the pathogenic role of myoglobin in rhabdomyolysis-induced acute kidney injury must rhabdomyolysis-induced acute kidney injury, pre kalemia, which occurs very early in the course of ied. Although plasmapheresis has been shown extracellular to the intracellular space (e.g., hyper venous hemofiltration or hemodiafiltration has temporarily, and the only means of removing principally with the use of super high-flux filters liuresis), the use of intestinal potassium binders, However, the evidence is mainly from isolated should not be treated unless it is symptomatic or known. In addition, some studies have shown unless severe hyperkalemia is present. Calcium- that the half-life of serum myoglobin does not to treat hyperphosphatemia, since the calcium load treated conservatively and those who receive con randomized studies are performed, preventive

The use of antioxidants and free-radical scav

engers (e.g., pentoxifylline, vitamin E, and vita min C) may be justified in the treatment or pre vention of myoglobinuric acute kidney injury, 52 as suggested by small case series, case reports, and various experimental studies of myoglobinuria, but controlled studies evaluating their efficacy are lacking.

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REFERENCES

- 1. Tein I, DiMauro S, Rowland LP. Myoglobinuria. In: Rowland LP, DiMauro S, eds. Myopathies. Handbook of clinical neurology. Vol. 62. Amsterdam: Elsevier Science Publishers, 1992:553-93.
- **2.** Warren JD, Blumbergs PC, Thompson PD. Rhabdomyolysis: a review. Muscle Nerve 2002;25:332-47.
- **3.** Allison RC, Bedsole DL. The other medical causes of rhabdomyolysis. Am J Med Sci 2003:326:79-88.
- **4.** Bagley WH, Yang H, Shah KH. Rhabdomyolysis. Intern Emerg Med 2007;2: 210-8
- **5.** Fernandez-Sola J, Grau JM, Pedro-Botet JC, et al. Nontraumatic rhabdomy-olysis: a clinical and morphological analy sis of 53 cases. Med Clin (Barc) 1988;90: 199-202. (In Spanish.)
- **6.** Giannoglou GD, Chatzizisis YS, Misirli G. The syndrome of rhabdomyolysis: pathophysiology and diagnosis. Eur J Intern Med 2007;18:90-100.
- 7. Wrogemann K, Pena SD. Mitochondrial calcium overload: a general mechanism for cell-necrosis in muscle diseases. Lancet 1976;1:672-4.
- **8.** Vanholder R, Sever MS, Erek E, Lameire N. Rhabdomyolysis. J Am Soc Nephrol 2000;11:1553-61.
- 9. Holt SG, Moore KP. Pathogenesis and treatment of renal dysfunction in rhab-domyolysis. Intensive Care Med 2001;27: 803-11.
- **10.** Melli G, Chaudhry V, Cornblath DR. Rhabdomyolysis: an evaluation of 475 hos pitalized patients. Medicine (Baltimore) 2005;84:377-85.
- 11. Ward MM. Factors predictive of acute renal failure in rhabdomyolysis. Arch Intern Med 1988:148:1553-7.
- 12. Veenstra J, Smit WM, Krediet RT, Arisz L. Relationship between elevated cre atine phosphokinase and the clinical spee trum of rhabdomyolysis. Nephrol Dial Transplant 1994:9:637-41.
- **13.** de Meijer AR, Fikkers BG, de Keijzer MH, van Engelen BG, Drenth JP. Serum creatine kinase as predictor of clinical course in rhabdomyolysis: a 5-year intensive care survey. Intensive Care Med 2003; 29:1121-5.
- **14.** Woodrow G, Brownjohn AM, Turney JH. The clinical and biochemical features of acute renal failure due to rhabdomyolysis. Ren Fail 1995;17:467-74.

15. Knochel JP. Rhabdomyolysis and myo globinuria. Annu Rev Med 1982;33:435-43. **16.** Zager RA, Gamelin LM. Pathogenetic mechanisms in experimental hemoglobin uric acute renal failure. Am J Physiol

1989;256:F446-F455.

- 17. Zager RA. Studies of mechanisms and protective maneuvers in myoglobinuric acute renal injury. Lab Invest 1989;60:619-29.
- 18. Zager RA, Foerder CA. Effects of inorganic iron and myoglobin on in vitro proximal tubular lipid peroxidation and cytotoxicity. J Clin Invest 1992;89:989-95
- 19. Reeder BJ, Wilson MT. Hemoglobin and myoglobin associated oxidative stress: from molecular mechanisms to disease states. Curr Med Chem 2005;12:2741-51.
 20. Holt S, Reeder B, Wilson M, et al. Increased lipid peroxidation in patients with
- rhabdomyolysis. Lancet 1999;353:1241. **21.** Bonventre JV, Weinberg JM. Recent ad vances in the pathophysiology of ischemic acute renal failure. J Am Soc Nephrol 2003;14:2199-210.
- **22.** Gabow PA, Kaehny WD, Kelleher SP. The spectrum of rhabdomyolysis. Medicine (Baltimore) 1982;61:141-52.
- 23. Hatamizadeh P, Najafi I, Vanholder R, et al. Epidemiologic aspects of the Bam earthquake in Iran: the nephrologic perspective. Am J Kidney Dis 2006;47:428-20
- **24.** Sieb JP, Penn AS. Myoglobinuria. In: Engel AG, Franzini-Armstrong C, eds. Myology. New York: McGraw-Hill, 2004: 1677-92.
- **25.** Simerville JA, Maxted WC, Pahira JJ. Urinalysis: a comprehensive review. Am Fam Physician 2005;71:1153-62. [Erratum, Am Fam Physician 2006;74:1096.]
- **26.** Wakabayashi Y, Kikuno T, Ohwada T, Kikawada R. Rapid fall in blood myoglobin in massive rhabdomyolysis and acute renal failure. Intensive Care Med 1994;20: 109-12.
- 27. Mikkelsen TS, Toft P. Prognostic value, kinetics and effect of CVVHDF on serum of the myoglobin and creatine kinase in critically ill patients with rhabdomyolysis. Acta Anaesthesiol Scand 2005;49:859-64.
 28. Grossman RA, Hamilton RW, Morse BM, Penn AS, Goldberg M. Nontraumatic rhabdomyolysis and acute renal failure. N Engl J Med 1974;291:807-11.

- **29.** Oh MS. Does serum creatinine rise faster in rhabdomyolysis? Nephron 1993; 63:255-7.
- **30.** Corwin HL, Schreiber MJ, Fang LS. Low fractional excretion of sodium: occurrence with hemoglobinuric- and myoglobinuric-induced acute renal failure. Arch Intern Med 1984:144:981-2.
- **31.** Better OS, Stein JH. Early management of shock and prophylaxis of acute renal failure in traumatic rhabdomyolysis. N Engl J Med 1990;322:825-9.
- **32.** Akmal M, Bishop JE, Telfer N, Norman AW, Massry SG. Hypocalcemia and hypercalcemia in patients with rhabdomyolysis with and without acute renal failure. J Clin Endocrinol Metab 1986;63:137-42.
- **33.** Sever MS, Vanholder R, Lameire N. Management of crush-related injuries after disasters. N Engl J Med 2006;354:1052-63
- 34. Shimazu T, Yoshioka T, Nakata Y, et al. Fluid resuscitation and systemic complications in crush syndrome: 14 Hanshin-Awaji earthquake patients. J Trauma 1997; 42:641-6
- **35.** Gunal AI, Celiker H, Dogukan A, et al. Early and vigorous fluid resuscitation prevents acute renal failure in the crush victims of catastrophic earthquakes. J Am Soc Nephrol 2004;15:1862-7.
- **36.** Homsi E, Barreiro MF, Orlando JM, Higa EM. Prophylaxis of acute renal failure in patients with rhabdomyolysis. Ren Fail 1997:19:283-8.
- **37.** Brown CV, Rhee P, Chan L, Evans K, Demetriades D, Velmahos GC. Preventing renal failure in patients with rhabdomyoly sis: do bicarbonate and mannitol make a difference? J Trauma 2004;56:1191-6.
- **38.** Cho YS, Lim H, Kim SH. Comparison of lactated Ringer's solution and 0.9% sa line in the treatment of rhabdomyolysis in duced by doxylamine intoxication. Emerg Med J 2007;24:276-80.
- **39.** Ron D, Taitelman U, Michaelson M, Bar-Joseph G, Bursztein S, Better OS. Pre vention of acute renal failure in traumatic rhabdomyolysis. Arch Intern Med 1984; 144:277-80.
- **40.** Richards JR. Rhabdomyolysis and drugs of abuse. J Emerg Med 2000;19:
- **41.** Moore KP, Holt SG, Patel RP, et al. A causative role for redox cycling of myoglobin and its inhibition by alkalinization

- in the pathogenesis and treatment of rhab domyolysis-induced renal failure. J Biol Chem 1998;273:31731-7.
- 42. Heyman SN, Greenbaum R, Shina A, Rosen S, Brezis M. Myoglobinuric acute renal failure in the rat: a role for acidosis? Exp Nephrol 1997;5:210-6.
- 43. Ho AM, Karmakar MK, Contardi LH, Ng SS, Hewson JR. Excessive use of normal saline in managing traumatized patients in shock: a preventable contributor to acidosis. J Trauma 2001;51:173-7.
- 44. Zager RA, Foerder C, Bredl C. The influence of mannitol on myoglobinuric acute renal failure: functional, biochemi-

- Soc Nephrol 1991;2:848-55.
- 45. Better OS, Rubinstein I, Winaver JM, Knochel JP. Mannitol therapy revisited (1940-1997). Kidney Int 1997;52:886-94. 46. Visweswaran P, Massin EK, Dubose TD Jr. Mannitol-induced acute renal failure. J Am Soc Nephrol 1997;8:1028-33.
- 47. Lameire N, Van Biesen W, Vanholder R. Acute renal failure. Lancet 2005;365:417-
- 48. Kellum JA. The use of diuretics and dopamine in acute renal failure: a systematic review of the evidence. Crit Care 1997; 1:53-9.
- cal, and morphological assessments. J Am 49. Evans KJ, Greenberg A. Hyperkalemia: a review. J Intensive Care Med 2005;20:
 - 50. Szpirt WM. Plasmapheresis is not justified in treatment of rhabdomyolysis and acute renal failure. J Cardiovasc Surg (Torino) 1997;38:557.
 - 51. Ronco C. Extracorporeal therapies in acute rhabdomyolysis and myoglobin clear ance. Crit Care 2005;9:141-2.
 - 52. Huerta-Alardin AL, Varon J, Marik PE. Bench-to-bedside review: rhabdomyolysis - an overview for clinicians. Crit Care 2005:9:158-69.

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