Hyperkalemic Emergency: When You Have Taken a Few Too Many KCl Tablets

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Abstract

Hyperkalemia is a common clinical problem that varies significantly in severity and indications for treatment. Hyperkalemic emergency exists when there are clinical signs or symptoms, including cardiac conduction abnormalities. The combination of nebulized albuterol and insulin with glucose is most effective for managing clinically significant hyperkalemia. Prompt recognition of hyperkalemic emergency, immediate interventions to lower extracellular potassium, and involvement of multiple disciplines (including critical care and nephrology) are essential to addressing this life-threatening presentation.

Categories: Emergency Medicine, Internal Medicine, Medical Education
Keywords: hyperkalemia, hyperkalemic emergency, potassium, critical care, emergency medicine, internal medicine, ekg, hemodialysis, insulin, bipolar disorder

Introduction

Hyperkalemia is a common clinical problem that presents in both the emergency and inpatient medicine settings. The frequency of hyperkalemia in the United States is reported to be between 2.6% and 3.2% in the general population [1]. Potassium is one of the most abundant cations in body [2], and disturbances to its homeostasis can have serious and even fatal consequences. Several factors can precipitate the formation of hyperkalemia, such as poor renal function, the use of certain medications, and independent demographic features [3,4]. The severity of hyperkalemia drives its management, which may be primarily through medical or procedural techniques, or through a combination of the two [1-15]. In this case report, we present a case of severe hyperkalemia due to combination of medication and supplement overdose, which required emergent hemodialysis due to hemodynamic instability.

Case Presentation

A 48-year-old female with a history of bipolar disorder, hypertension, and gastro-esophageal reflux disease (GERD) presented to the emergency department stating, "I have low potassium." She reported having hypokalemia for the past 10 years and had been taking 60 mEq daily of supplemental potassium chloride (KCl). She began having muscle cramps one day prior to presentation and decided to "double up" on her KCl tablets, thinking her symptoms were due to hypokalemia. She denied any recent depressive symptoms or suicidal ideation. She had two days of non-bloody, non-bilious vomiting associated with multiple episodes of non-bloody diarrhea. She denied fevers, chills, or night sweats.

On presentation to the emergency department, vital signs were temperature 97.1 degrees Fahrenheit, heart rate 52 beats per minute, blood pressure 135/74 mmHg, and respiratory rate 17 breaths per minute. On physical examination, she was awake, alert, and oriented. She had a normal physical exam including focused cardiovascular and neurological examinations. The most remarkable abnormal laboratory results were serum potassium (K) 9.6 mmol/L, serum chloride (Cl) 117 mmol/L, blood urea nitrogen (BUN) 73 mg/dL, and creatinine 4.67 mg/dL (with no known baseline). White blood cell (WBC) count was elevated at 13.28 K/mm³ with normal differential. All laboratory values are shown in Table 1.
### Laboratory Test Results

| Laboratory Test                     | Patient’s Result | Reference Range          |
|------------------------------------|------------------|--------------------------|
| White blood cell count (WBC)       | 13.28 K/mm³³      | 4.0-12.0 K/mm³³          |
| Hemoglobin (Hb)                    | 12.5 g/dL        | 12.0-16.0 g/dL          |
| Platelet count (Plt)               | 306 K/mm³³       | 130-400 K/mm³³          |
| Sodium (Na)                        | 136 mmol/L       | 136-145 mmol/L          |
| Potassium (K)                      | 9.6 mmol/L       | 3.7-5.1 mmol/L          |
| Chloride (Cl)                      | 117 mmol/L       | 98-107 mmol/L           |
| Bicarbonate (CO₂)                  | 16 mmol/L        | 21-32 mmol/L            |
| Blood urea nitrogen (BUN)          | 73 mg/dL         | 7-18 mg/dL              |
| Creatinine                         | 4.67 mg/dL       | 0.55-1.3 mg/dL          |
| Glucose                            | 101 mg/dL        | 74-106 mg/dL            |
| Calcium                            | 9.4 mg/dL        | 8.4-10.1 mg/dL          |
| Phosphorus                          | 3.3 mg/dL        | 2.5-4.9 mg/dL           |

**TABLE 1: Initial Laboratory Results**

Initial electrocardiogram (ECG) showed normal sinus rhythm with frequent premature ventricular contractions in a pattern of bigeminy (Figure 1). Repeat electrocardiogram showed sinus bradycardia with heart rate (HR) 47 bpm and peaked T waves (Figure 2).

**FIGURE 1: Initial Electrocardiogram (ECG)**

This is the initial ECG upon the patient's arrival to the emergency department. It shows normal sinus rhythm with frequent premature ventricular contractions in the pattern of bigeminy. Peaked T waves are labeled with arrows.
The patient received treatment with 50 mEq IV sodium bicarbonate, 1 gram IV calcium gluconate, five units IV insulin with one ampule of D50, and polysterene sulfate 60 grams orally. Shortly after, she became progressively bradycardic and hypotensive. An emergent trialysis catheter was placed, and she was started on a norepinephrine drip and began emergent hemodialysis.

Her serum potassium normalized after hemodialysis to 4.2 mmol/L and her creatinine improved to 1.29 mg/dL at discharge. The patient was evaluated by the psychiatry team due to concerns for self-harm; however, the patient denied, and her ingestion was ultimately determined to be a therapeutic misadventure.

Discussion
As a primarily intracellular cation, potassium has an inherent concentration gradient between its intracellular and extracellular components [1,2,13,14]. This concentration gradient is the basis of the cellular resting membrane potential, and as such, potassium homeostasis is the basis for cellular integrity. Hyperkalemia is defined as a serum potassium >5.0 mmol/L, and can be the result of increased potassium absorption, decreased potassium excretion, or a shift of potassium across the cell membrane [2,14].

Hyperkalemia contributes to cardiac arrhythmias by destabilizing the myocardial conduction system [12]. Decreased resting membrane potential in the setting of hyperkalemia leads to increased cardiac depolarization, myocardial excitation, and cardiac instability, with subsequent arrhythmia formation including ventricular fibrillation and asystole. Tall, peaked T waves are typically the earliest manifestations of hyperkalemia and occur before changes to the QRS complex. In the absence of any other ECG findings, peaked T waves are rarely associated with life-threatening arrhythmias. QRS complex changes (uniform widening) are often evident with increasingly severe hyperkalemia. Loss of P waves then follows, ultimately leading to the formation of sine waves - the fusion of wide QRS complexes with ST-T segments. Additionally, severe hyperkalemia can manifest with sinus bradycardia or arrest, atrioventricular (AV) blocks, and loss of pacemaker capture. Hyperkalemia can impair conduction in the His-Purkinje system and can also cause fascicular and bundle branch blocks.

IV insulin, often along with one ampule of D50 to avoid hypoglycemia, is given to lower the serum potassium concentration by driving potassium intracellularly and enhancing the activity of the Na/K ATPase pump, thereby promoting intracellular potassium shifts, taking effect within 10-15 minutes. Insulin should not be given to patients who have concurrent basal or long-acting insulin, including subcutaneous infusion pumps. Albuterol 10-15 mg nebulized is also used to transfer potassium intracellularly, with onset of 60-90 minutes. The combination of insulin/glucose and nebulized albuterol are likely most effective for lowering potassium emergently [16-18].

Calcium gluconate is commonly used in the management of life-threatening arrhythmias in the setting of hyperkalemia due to immediate onset and effects. Adult dosing is 1.5-3 g IV q2-5 minutes PRN with a maximum dosing of 200 mg/min or 3 g/episode or 15 g/day. Adverse reactions include hypercalcemia, accompanied by vasodilation, hypotension, bradycardia, syncope, and dizziness, as well as more severe symptoms such as subsequent arrhythmias and extravasation necrosis. Sodium bicarbonate also works by transcellular potassium shifting, but has limited demonstrated efficacy in lower serum K levels. Sodium polystyrene sulfonate (Kayexalate) has been used in the management of hyperkalemia as a potassium
Hyperkalemia is a common clinical problem that varies significantly in severity and indications for treatment. Hyperkalemia in the emergent setting is when there are clinical signs or symptoms including conduction abnormalities or muscle paralysis. The combination of nebulized albuterol and insulin with glucose is most effective for managing clinically significant hyperkalemia. Intravenous calcium will only stabilize the cardiac membrane for around 30-60 minutes and is a transient measure that should be used as an adjunct with definitive serum potassium-lowering treatments. Prompt recognition of hyperkalemic emergency and involvement of multiple disciplines (including critical care and nephrology) is essential to addressing this life-threatening presentation.

Conclusions

Hyperkalemia is a common clinical problem that varies significantly in severity and indications for treatment. Hyperkalemic emergency exists when there are clinical signs or symptoms including conduction abnormalities or muscle paralysis. The combination of nebulized albuterol and insulin with glucose is most effective for managing clinically significant hyperkalemia. Intravenous calcium will only stabilize the cardiac membrane for around 30-60 minutes and is a transient measure that should be used as an adjunct with definitive serum potassium-lowering treatments. Prompt recognition of hyperkalemic emergency and involvement of multiple disciplines (including critical care and nephrology) is essential to addressing this life-threatening presentation.

Additional Information

Disclosures

Human subjects: Consent was obtained by all participants in this study. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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