

Death Following Recreational Use of Designer Drug “Bath Salts” Containing 3,4-Methylenedioxypyrovalerone (MDPV)

Brittany L. Murray · Christine M. Murphy ·
Michael C. Beuhler

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Abstract

Introduction 3,4-Methylenedioxypyrovalerone (MDPV) is a designer stimulant drug that has gained popularity in the USA. Although adverse effects of MDPV have been described, to our knowledge, this is the first reported death.

Case Report We report the case of a 40-year-old male who injected and snorted “bath salts” containing MDPV and subsequently became agitated, aggressive, and experienced a cardiac arrest. He was resuscitated after his initial arrest; however, he developed hyperthermia, rhabdomyolysis, coagulopathy, acidosis, anoxic brain injury, and subsequently died.

Discussion This is the first case in the medical literature to report death due to isolated confirmed MDPV intoxication. The manner of death is also consistent with excited delirium syndrome.

Keywords 3,4-Methylenedioxypyrovalerone · Bath salts · Excited delirium syndrome · MDPV · Designer drug

Introduction

Psychoactive products containing β -keto phenylalkylamines such as 3,4-methylenedioxypyrovalerone (MDPV), mephedrone, and methylone have entered the recreational

drug market over the past several years. These products are often labeled as “bath salts” or “plant food,” with the disclaimer that they are “not for human consumption” to skirt state and federal laws. Although states are creating new laws to address these derivatives, an increasing number of exposures have been reported to United States (US) poison centers since late 2010 [1].

MDPV is a ring-substituted analogue of pyrovalerone (Fig. 1). Pyrovalerone is a stimulant and a schedule V controlled substance that was first synthesized in 1964. The synthesis of MDPV was first reported in 1969 [2, 3]. The chemical structure of MDPV is similar to methcathinone (“Mcat”) and to hallucinogenic substances like 3,4-methylenedioxymethamphetamine (MDMA or “Ecstasy”), but it is best characterized as a β -keto phenylalkylamine (Fig. 1). Over the past decade, MDPV has gained popularity as a designer drug, or “legal high,” across Europe. The first designer drug containing MDPV was identified in Germany in 2007 [4]. In Japan, MDPV was retrospectively identified in designer drugs confiscated in the year 2006 [5]. The recreational use of MDPV in the USA has become more prevalent since late 2010 and it is now illegal in many states [1]. We report the first case of isolated recreational use of MDPV resulting in excited delirium syndrome and ultimately death, with confirmatory toxicological analyses.

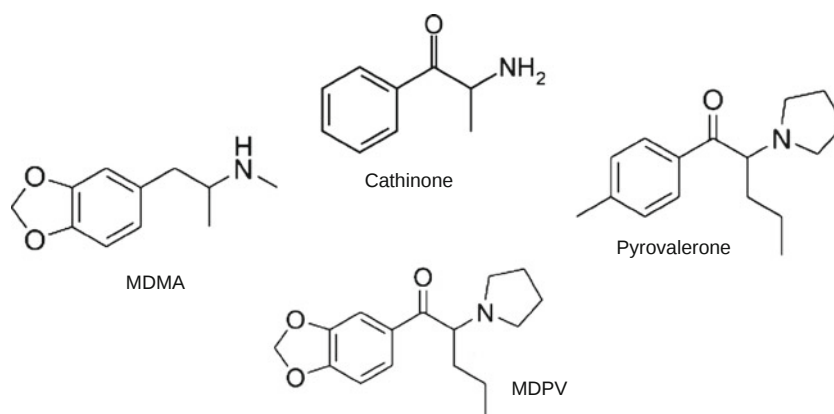
Case Report

A 40-year-old man with a history of bipolar disorder snorted and injected an unknown amount of “bath salts.” Family and friends reported he had recently switched from abusing cocaine to using “bath salts” products. Shortly after his consumption of this product, he became aggressive, uncontrollable, delusional, removed all of his clothing, and

B. L. Murray · C. M. Murphy (✉)
Department of Emergency Medicine, Carolinas Medical Center,
PO Box 32861, MEB 3rd Floor,
Charlotte, NC 28232, USA
e-mail: christine.murphy66@gmail.com

M. C. Beuhler
Carolinas Poison Center,
Charlotte, NC, USA

Fig. 1 Chemical structures of 3,4-methylenedioxypropylvalerone (MDPV); cathinone; 3,4-methylenedioxyamphetamine (MDMA); and pyrovalerone

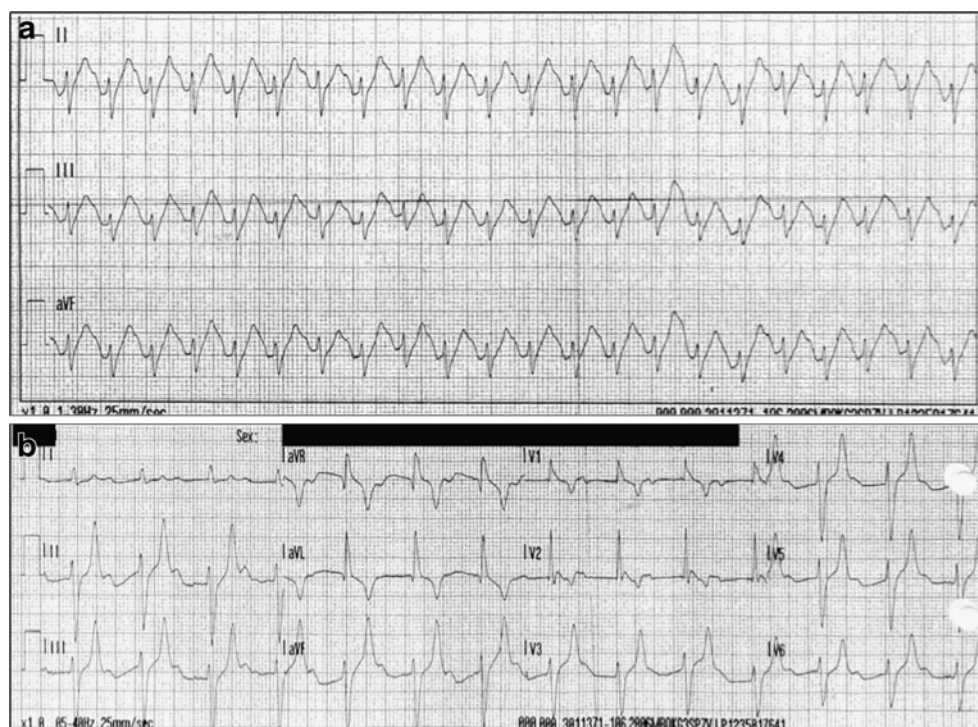


ran outside. Police were called and while being taken into custody, the patient displayed aggression, considerable strength, and violent behavior. An electronic control device was discharged three times in an effort to overpower him and protect others on scene. During ambulance transport, he remained aggressive and delusional and was physically restrained. He was noted to have slightly labored breathing and was placed on a non-rebreather mask (NRB) with 100% oxygen. He was yelling incomprehensibly and was noted to have dilated pupils. Initial vital signs in the prehospital setting were as follows: heart rate 164 beats per minute (bpm); blood pressure 131/72 mmHg; respiratory rate 24 breaths/min, and oxygen saturation of 100% on NRB. Prehospital electrocardiogram (EKG) initially demonstrated sinus tachycardia with widened QTc interval and peaked T waves (Fig. 2a). Repeat EKG 10 min later depicted

normal sinus rhythm with persistent peaked T waves (Fig. 2b). Sedation was attempted unsuccessfully with 2 mg of intramuscular lorazepam.

Upon arrival in the hospital, he continued with very aggressive behavior and incomprehensible screaming. A review of the patient's electronic medical records revealed previous routine medications of quetiapine, methadone, temazepam, and 10/650 mg hydrocodone/acetaminophen. As the patient was never conversant, compliance was not established. Vital signs at the time of his arrival, 15 min after reported EMS vital signs, were as follows: oral temperature 98.0°F; blood pressure 100/64 mmHg; heart rate 91 bpm; respiratory rate 12 breaths/min; and oxygen saturation of 100% on NRB. While being transitioned from the Emergency Medical Services stretcher to a hospital bed, and without further intervention, he became very quiet and withdrawn.

Fig. 2 **a** Initial prehospital EKG. **b** Repeat prehospital EKG performed 10 min later demonstrating normal sinus rhythm with rate of 85 bpm, PR interval 110 msec, QRS interval 116 msec, QTc interval 414 msec, and peaked T waves



Within 5 minutes of his arrival to the hospital he developed bradycardia and subsequent cardiac arrest with pulseless electrical activity (PEA). Standard advanced cardiac life support measures were initiated including cardiopulmonary resuscitation and administration of two doses each of 1 mg intravenous (IV) epinephrine and 1 mg IV atropine, as well as 100 mg of IV lidocaine. He was intubated and given both 2 mg IV naloxone and 0.5 mg IV flumazenil without effect. Return of spontaneous circulation was achieved after 30 min of resuscitation. Dopamine and phenylephrine infusions were initiated at rates of 5 mcg/kg/h and 10 mcg/h, respectively, due to persistent hypotension despite a bolus infusion of 2 liters 0.9% normal saline (NS).

Immediately after resuscitation, a rectal temperature was 105.4°F, for which he was given 1,300 mg of acetaminophen per rectum. Other vital signs included a blood pressure of 70/32 mmHg; heart rate of 91 bpm; and oxygen saturation of 100% while mechanically ventilated. Physical examination revealed a Glasgow coma scale score of 3 and pupils dilated at 6 mm with minimal reaction to light. Skin examination revealed dry skin with multiple needle marks in both antecubital fossae. Additionally, there was no evidence of hypertonia, hyperreflexia, inducible or spontaneous clonus, or bruxism.

Pertinent initial labs in the immediate post-arrest period included: potassium of 7.4 mmol/L, creatinine of 3.0 mg/dL, negative serum acetaminophen and ethanol levels (limit of detection less than or equal to 5 mg/dL), a salicylate level of 4.1 mg/dL, and a urine drug screen positive for opiates but negative for cocaine, phencyclidine, amphetamine, tetrahydrocannabinol, benzodiazepines, and barbiturates. Initial international normalized ratio (INR) was 1.01; creatinine

kinase 234 U/L; aspartate aminotransferase (AST) 19 U/L; and alanine transaminase (ALT) was 36 U/L. An EKG was also obtained at this time and showed changes consistent with hyperkalemia (peaked T waves) with a sinus bradycardia at a rate of 56 bpm with a markedly prolonged QRS of 240 milliseconds (msec) (Fig. 3). The patient was then transferred to a tertiary hospital with hemodialysis capabilities and a toxicology consult service.

On arrival to the tertiary care center, the patient's temperature had decreased to 100.2°F without further intervention. The patient's blood pressure remained low at 85/41 mmHg despite elevated doses of dopamine (20 mcg/kg/min), phenylephrine (180 mcg/h), and additional boluses of chilled IV NS (4 liters); therefore, norepinephrine was started (4 mcg/min). His heart rate was 115 bpm and oxygen saturation of 100% with the patient breathing over the set ventilator rate of 20 breaths/min at 32 breaths/min. His pupils remained dilated and were minimally reactive to light; however, on neurologic examination, he had a normal gag reflex, normal corneal reflexes, and he flexed in response to pain with all four extremities. Repeat EKG showed a bradycardia with a rate of 53 bpm, a right bundle branch block with hyperacute T waves, some ST depression in V2 and V3, a QRS interval of 158 msec, and QTc interval of 420 msec. A venous blood gas was performed: pH 7.2, pCO₂ 39 mmHg, pO₂ 35 mmHg, HCO₃ 16.2 mmol/L, base excess -11 mEq/L, and serum lactate of 2.83 mmol/L. Laboratory testing repeated 5 h after initial presentation demonstrated continued hyperkalemia (8.0 mmol/L), an elevated creatinine (3.5 mg/dL), and marked increase in AST (869 U/L), ALT (738 U/L), INR (4.2), and creatinine kinase (14,839 U/L). His hyperkalemia was treated with 1 g of calcium gluconate, 50 mEq of sodium

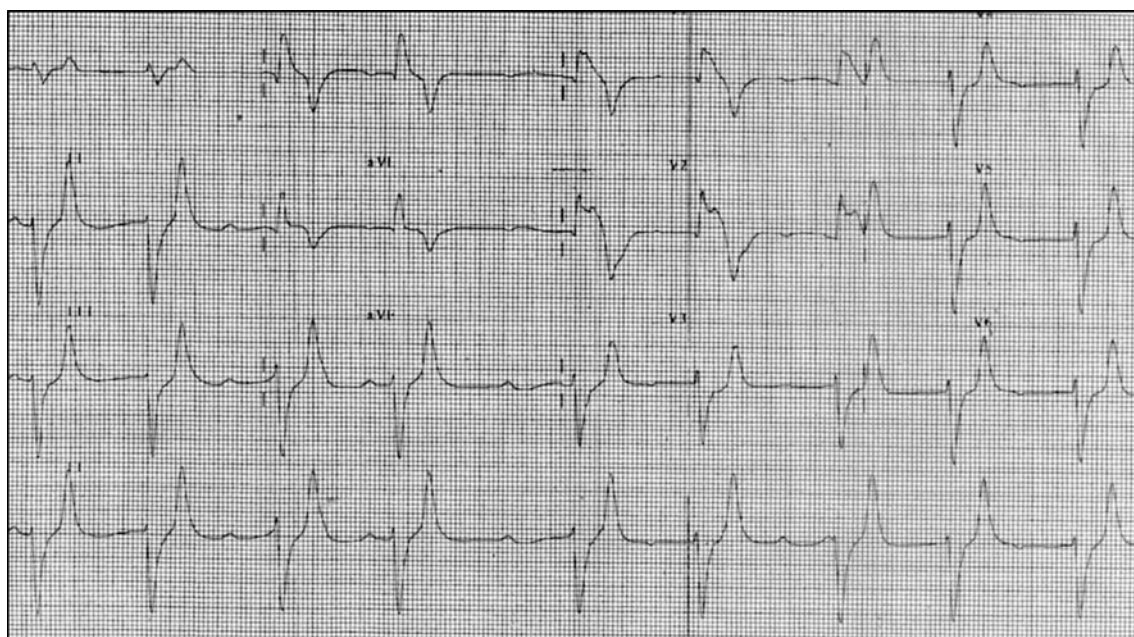


Fig. 3 Post-arrest EKG

bicarbonate, 10 units of regular insulin, and 50 g of dextrose. His potassium subsequently decreased to 6.0 mmol/L and a repeat EKG demonstrated sinus tachycardia (120 bpm) with normalization of the T waves (Fig. 4).

He was admitted to the medical intensive care unit where he was started on an infusion of 5% dextrose in water with 150 mEq of sodium bicarbonate at a rate of 125 ml/h, hydrocortisone (100 mg IV every 8 h), and propofol infusion (10 mcg/kg/h). Persistent hypotension was treated with an additional 2 liters of bolused NS and a vasopressin infusion was started at 0.04 units/min. Over the next 3 h, he developed a worsening metabolic acidosis (arterial pH 7.14; HCO_3^- 10 mmol/L) with oliguric renal failure that progressed to anuria (creatinine 4.26 mg/dL), worsening transaminase elevation (AST 10,873 U/L; ALT 6,623 U/L), and rhabdomyolysis (creatinine kinase 75,952 U/L). His coagulopathy progressed and his INR was >9.3 (upper limit of laboratory detection) and PT >96 s (upper limit of laboratory detection). He was noted to be very pale and began having melanotic stools. He developed an anemia (hemoglobin of 6.3 g/dL) and thrombocytopenia with platelets of $11 \times 10^9/\text{L}$. Hemodialysis was started 17 h after presentation to correct the acidosis and renal failure. Additionally, he received packed red blood cells (4 units), platelets (2 units), cryoprecipitate (9 units), and fresh frozen plasma (10 units). A non-contrasted brain computed tomography showed decreased gray–white matter discrimination interpreted by the radiologist as “likely edema and early anoxic injury.” An EEG was performed and showed diffuse slowing consistent with coma and likely anoxic injury.

Despite improvement in his acidosis and anemia after dialysis and transfusion, his neurologic exam worsened. His pupils became fixed and dilated and he lost his

gag, corneal, and vestibulo-ocular reflexes. He no longer responded to noxious stimuli. Approximately 42 h after his initial presentation, the patient was declared brain dead by clinical criteria and support was withdrawn.

During his evaluation, several toxicology laboratories were evaluated on blood and urine samples obtained at the time of arrival to the tertiary care center. His urine was negative for barbiturates, amphetamines, benzodiazepines, cocaine, marijuana, methadone, and opiates. Urine liquid chromatography was positive for trimethoprim. Serum ethanol levels were 11 mg/dL (8 h after presentation) and 25 mg/dL (19 h after presentation). His qualitative urine methadone screen was negative. Serum salicylate level was 7.9 mg/dL and serum acetaminophen level was 2.9 mcg/mL. A lithium level was <0.1 mmol/L. Ethanol, ethylene glycol, methanol, and isopropanol were not detected by gas chromatography 21 h after presentation.

Samples of the patient’s urine and serum obtained at the time of his arrival to the tertiary care center were sent for further testing at NMS Labs (Forensic Science Department, Willow Grove, PA). An initial therapeutic drug screen using gas chromatography/mass spectrometry was performed and positive for acetaminophen, caffeine, cotinine, lidocaine, trimethoprim, and MDPV; quetiapine was not detected. Confirmatory testing was undertaken using high performance liquid chromatography/tandem mass spectrometry (LC/MSMS). Specimens were fortified with d_8 -MDPV as an internal standard and then subjected to a single step liquid-liquid extraction using trichloroacetic acid (TCA). MDPV was quantified in his urine (670 ng/mL) and serum (82 ng/mL). Trimethoprim was also quantified using high performance thin layer chromatography (12 mcg/mL in urine and 2.2 mcg/mL in serum).

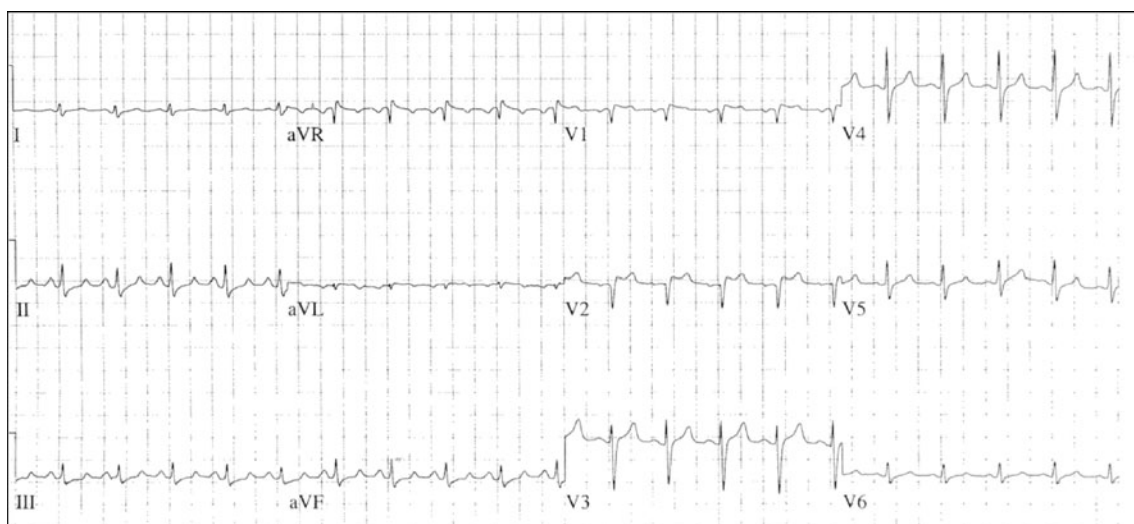


Fig. 4 EKG after treatment for hyperkalemia. PR interval of 125 msec, QRS interval of 83 msec, QTc interval of 412 msec, normalization of the T waves and ST depressions in leads II and III with mild ST elevation in leads V1 and V2

Discussion

MDPV is a white or tan powder that is insufflated, injected intravenously, or ingested. Although few studies have formally examined the clinical effects or metabolism of MDPV in humans, multiple first-hand reports and a handful of published case reports now exist [6–10]. Stimulant effects, an increase in libido, anxiety, hallucinations, and paranoid psychosis have been reported [6]. MDPV appears to undergo CYP450 2D6, 2C19, and COMT phase 1 metabolism into methylcatechol and pyrrolidine, which in turn are glucuronidated to allow for renal excretion [11, 12]. Although the mechanism of action is not fully elucidated at this time, a dopaminergic mechanism is hypothesized due to the chemical similarity to amphetamines and the behavioral effects of MDPV that mimic those of amphetamines which are known to affect dopaminergic pathways. Studies on other pyrovalerone analogs demonstrated potent inhibition of the dopamine and norepinephrine transporters responsible for reuptake of their corresponding neurotransmitters [13]. Further supporting this hypothesis are animal studies demonstrating MDPV increases dopamine levels in experimental mice and rats [14, 15].

Since late 2010, MDPV has been the most commonly detected β -keto phenylalkylamine in toxicologic analysis of “bath salts,” “plant foods,” or other legal highs in the USA. This differs from the European experience where mephedrone was most commonly detected in these substances [1, 9, 16]. This difference has an unknown clinical significance, as reports of β -keto phenylalkylamine use in the USA describe similar sympathomimetic effects of both mephedrone and MDPV, including agitation, hypertension, palpitations, hallucinations, and violent behavior; however, the US reports have differed in part from those from Europe by the increased prevalence of MDPV, as well as an increase in cases involving rhabdomyolysis.

We report a case of MDPV toxicity with symptoms consistent with Excited Delirium Syndrome (ExDS) followed by a sudden PEA arrest, return of spontaneous circulation, and subsequent development of coagulopathy, rhabdomyolysis, renal failure, hepatic failure, anoxic brain injury, and death. This is the first case in the medical literature to report death due to isolated MDPV intoxication. Toxicological analysis of serum and urine revealed MDPV as an isolated intoxicant. Additionally, while a death consistent with ExDS has been reported after mephedrone use, to our knowledge, this is the first reported MDPV-associated death [17].

ExDS is characterized by delirium with agitation, hyperthermia, tachypnea, tachycardia, a period of “giving up,” or cessation of struggle, followed by cardiac

arrest, as was seen in our patient [18]. ExDS most likely results from dysregulation of dopaminergic pathways. ExDS autopsy data have shown a decreased number of D3 dopamine receptors when compared to controls [19]. This suggests that ExDS patients may lack normal compensatory measures for dealing with rapid changes in dopamine levels. Furthermore, hypothalamic dopamine receptors are responsible for thermoregulation, and patients with ExDS have elevated heat shock proteins consistent with the clinically observed hyperthermia [19]. Hyperthermia in these patients is likely a sign of autonomic dysfunction that results in death in many ExDS patients. However, in patients that survive the initial arrest, such as the patient in this report, hyperthermia likely contributes to the coagulopathy, rhabdomyolysis, and multisystem organ failure that result in death after the initial insult. ExDS has been described with MDMA, cocaine, and amphetamine intoxication [20, 21].

In our case, the patient’s previous history of cocaine abuse may have also predisposed him to ExDS, as chronic cocaine abuse is known to alter neurotransmitters such as dopamine [22]. The dopaminergic effects of MDPV likely contributed to aberrant thermoregulation, and the sympathomimetic effects contributed to the agitation, delirium, and the cardiac arrest in this patient. Treatment for MDPV intoxication, along with other drugs in the stimulant class and patients with ExDS remains largely supportive, with the use of intravenous benzodiazepines, intravenous fluids, and cooling.

The patient in this case was also found to have trimethoprim present on serum and urine testing. This was quantified (12 mcg/mL in urine and 2.2 mcg/mL in serum) and present in amounts that exceed previous reports of peak plasma levels in single one-time ingestions of 160 mg doses [23]. Trimethoprim is not reported to cause stimulant effects with chronic use or in acute overdose. There are reports of hyperkalemia occurring in patients predisposed to potassium retention (underlying renal failure or concurrent diuretic use) [24, 25]. Our patient was not known to be taking a trimethoprim-containing medication, and personal communication with the laboratory performing the test (telephone correspondence with Dr. S. Kacinko, NMS Labs, August 2011) revealed an additional case where MDPV and trimethoprim were found together using similar testing methods. We hypothesize that this may be an adulterant in some formulations of “bath salts,” including that which our patient consumed. It is therefore possible trimethoprim contributed to our patient’s hyperkalemia, but we do not feel it is responsible for his ExDS symptoms or ultimate death.

We were initially perplexed as to why our patient’s serum ethanol level continued to rise well after admission even with the initiation of hemodialysis; however, ethanol

was not detected on a volatile alcohol screen performed on blood obtained during this time period. Most likely the serum ethanol levels quantified via the serum ethanol assay (Beckman UniCel Dx C, Beckman Coulter Inc, Brea, CA) were falsely elevated due to interference from the patient's profoundly elevated lactate dehydrogenase (LDH) (11,108 IU/L, 19 h after presentation) and lactate levels (10.7 mmol/L, 19 h after presentation). This interference of LDH and lactate with serum ethanol determination has been described with multiple instruments/assays, including the methodology used to measure ethanol in this case [26–28]. These assays are based on the conversion of ethanol to acetaldehyde by alcohol dehydrogenase reducing NAD to NADH in the process, which is quantitated by absorbance at 340 nm due to NADH formation. Assay interference occurs when excess LDH and lactate allow for conversion of NAD to NADH in a reaction unrelated to the presence of ethanol.

Conclusion

Recreational abuse of “bath salts” products continues despite legislative initiatives to prohibit its use. While this is the first reported case of death due to isolated recreational use of MDPV with confirmatory toxicological analyses, further characterization of symptoms related to intoxication and associated deaths is needed. This case highlights the need for emergency physicians and toxicologists to be vigilant in testing for novel drugs of abuse, as it allows for an improved understanding of the potential toxicity of these drugs and associated adulterants.

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