Two cases of paradoxical hypothermia in a Sahara desert multi-stage ultramarathon

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Abstract

Exertional hypothermia in a hot environment, a seemingly contradictory state, is a clinical presentation that has not been previously described in the medical literature. We present two cases of symptomatic hypothermia of 34.35°C (95°F) which resolved with ingestion of calories in otherwise healthy men competing in a multi-stage ultramarathon in the Sahara desert, Egypt 2012 with ambient temperatures of approximately 38-39°C (100-102.2°F). Hypothermia is well documented in the medical literature as a presenting sign of hypoglycemia in both conscious and comatose patients. The mechanism by which hypoglycemia contributes to hypothermia is poorly understood, but may represent a compensatory response that reflects a decrease in energy demand during glucose deprivation. Wilderness medicine practitioners at endurance events should be cognizant of hypothermia as a potential presenting sign of hypoglycemia, as it can be empirically treated and rapidly reversed.

Introduction

An ultramarathon is any race longer than the standard 42.2 km (26.2 mile) marathon.1 These races are gaining popularity in North America, with an almost 10% increase in annual participants, and more than 69,000 finishers in 2013.2 Competitors often run through a variety of terrain in inclement conditions with temperatures ranging from below freezing to greater than 40°C (104°F).3 Although the majority of medical injuries encountered are often minor dermatologic and musculoskeletal issues,4,5 other serious and potentially debilitating conditions like exercise-associated hyponatremia (EAH)6 or acute kidney injury7 have been reported as well.

Heat-related illnesses are a concern for any medical provider of ultramarathon runners with presentations that range from mild, usually self-limited heat exhaustion to potentially fatal heat stroke.7 Heat-related illness can present similarly to EAH8 or altitude illness,9 but a concomitant concern for, or presentation of hypothermia has not been discussed in recent practice guidelines.10 Exertional hypothermia was first documented in the temperature 1901 Boston Marathon in one runner who had consumed large amounts of alcohol, and in another who was exhausted more than the average.11 However, as there are no reported cases of exertional hypothermia in hot conditions, a seemingly contradictory state, we introduce and define the term, paradoxical hypothermia. We present two cases of hypothermia in otherwise healthy men competing in a multi-stage ultramarathon in the Sahara desert, Egypt with ambient temperatures above 38°C (100.4°F).

Case Report

Case #1

A 42 year old male with no significant past medical history and taking no medications was a participant in RacingThePlanet© 250 km (155 mile) 7 day, 6 stage ultramarathon in the Sahara desert, Egypt October, 2012. On the third stage the individual was observed at checkpoint 2, 15.1 km (9.4 miles) from the start with ambient temperatures of approximately 39°C (102.4°F), no wind or rain. He appeared fatigued with a pasty pale color, was walking slowly, and stated that he felt unwell and nauseous. He had been unable to tolerate his regular amount of water or electrolytes, and had resting heart rate of 84 beats per minute by palpitation. After 30 min of rest in the shade, the competitor continued on the race. On arrival at checkpoint 3, 23.5 km (14.7 miles) from the start, he continued to complain of fatigue and nausea. He was pale, with cool and diaphoretic skin, and complained of feeling cold. He had a resting heart of 80 beats per minute by palpitation, blood pressure of 116/75 mm Hg by manual monitor, with an axillary temperature of 34.4°C (94°F) (digital thermometer; The Kroger Co., Atlanta, GA, USA). He was provided with a 12oz soda (150 cal; Pepsi Co., Sunnyvale, CA, USA) and placed in a support vehicle to rest. After ingesting the soda his symptoms of coldness, nausea and fatigue resolved, and he continued on the course. The participant ultimately finished the stage.

Case #2

A 50 year old male with no significant past medical history and not taking any medications was a participant in the same Racing ThePlanet© event. The patient was seen on stage 2 at checkpoint 1 at 9 am 11.2 km (7.0 miles) from the start with an ambient temperature of approximately 38°C (100.4°F), with no wind or rain. He complained of feeling tired and had required frequent stops on the course, but felt better and continued to race after resting in the shade of the checkpoint. Further on the course he was seen several times hunched over, fatigued and apparently trying to catch his breath. He was stopped by a member of the medical team at 17 km (10.5 miles) from the start. He complained of feeling cold, with cool and moist skin. He appeared exhausted, without ataxia, and was leaning on the vehicle for support. He appeared agitated, repeatedly stating, “I’m OK, I can go on,” and resisted efforts of medical assistance. His heart rate was 86 beats per min by palpitation, respiratory rate of 20 breaths per min, with an oral temperature of 35.1°C (95.3°F) (digital thermometer; The Kroger Co.). A repeat temperature several minutes later was 35.0°C (95.0°F). The medical team member used the same thermometer to check his own oral temperature, which was 36.2°C (97.1°F). The competitor was withdrawn from the race, and symptoms of cold and fatigue resolved with ingestion of food and rest.
Discussion

There are little data that examine the incidence of hypothermia in geographical regions with a warm climate. Hypothermia is the condition when the body’s temperature falls to 35°C (95°F) or below,14 that can present as a primary environmental disease or a complicating secondary state to a medical or traumatic condition. In hypothermic patients presenting to the emergency department, there is a 7% mortality with initial temperatures of mild hypothermia between 35 and 32.2°C, and a 23% mortality when less that 32.2°C.14 The primary effect of cooling is a decrease in metabolism, but the body’s compensatory mechanisms are fully functional in cases of mild hypothermia, with ataxia, dysarthria and apathy commonly present.15 Factors that may contribute to the development of hypothermia range from environmental conditions such as wind, low temperatures, and moisture; to medical comorbidities such as diabetes, hypothyroidism, or substance abuse.16 It is highly unlikely that these two participants in a multi-stage ultramarathon in the hot and dry conditions of the Sahara desert presented with primary hypothermia. Rather, it is reasonable that it was a secondary manifestation to a disease such as hypoglycemia, a causative relationship that has been well described in hospital settings, but not previously in healthy ultramarathon runners.

The presenting symptoms and signs of hypoglycemia are well known to physicians. They may include anxiety, a sense of ill health, with feelings of detachment or apathy. There is often diaphoresis, ataxia, and if more severe, encephalopathy that may progress to coma and death if uncorrected. Hypothermia as a presenting sign of hypoglycemia was first described in 196417 and has been most frequently described in diabetics,18 but also in those with varying oncologic processes, liver disease, and alcohol abuse with and without starvation states.19 One of the early investigations into the association between hypothermia and hypoglycemia noted that 53% of consecutive hypoglycemic patients had hypothermia vs 5% of encephalopathy of other etiologies.19

While the exact mechanism of hypothermia in hypoglycemia remains unclear, it is hypothesized that it may be due to increased heat dissipation from cutaneous vasodilation caused by disorder of the heat-regulating hypothalamus.19 Experimental evidence for this centrally mediated mechanism was initially described by Freinkel et al. in studies that injected a synthetic intracellular glucose inhibitor that produced hypoglycemia despite a serum euglycemic state. Healthy male volunteers’ temperatures decreased 1.1°C (3.4°F) below baseline for 6 h, indicating that hypothermia was triggered by intracellular glucopenia rather than by the availability of circulating glucose.20 Further animal studies found a 5-fold hypothermic response to injected centrally acting glucose inhibitor versus peripheral exposure,21 supporting the theory that thermoregulatory centers in the hypothalamus are thought to be both glucose and insulin sensitive.22

These studies suggest that hypothermia is a direct consequence of neuroglucopenia, and the decreased body temperature may be protective by preserving neuronal function while decreasing peripheral energy demands during times of glucose deprivation.11,12

Blood glucose levels in humans represent a balance between tissue uptake, hepatic glucose production – which in turn is dependent on glucose intake – glycogen reserves and gluconeogenesis from amino acids and their precursors.22 With exercise, skeletal muscles can increase their metabolic consumption by up to 20 times.23 The ultramarathon participants in these 2 cases had high energy consumption and may have developed hypoglycemia due to increased tissue uptake from exercise, insufficient caloric intake, insufficient glycogen reserves, or a combination of all three. The initial hypoglycemia is likely what caused decreased neuronal intracellular glucose levels, which led to a disruption of the thermoregulatory centers in the hypothalamus. The rapid reversal of symptoms with intake of calories supports the supposition that the patients were hypoglycemic, as well as the proposed mechanism by which these athletes became hypothermic despite running through the heat of the Sahara desert.

Some limitations in this case report exist, namely the lack of measurement of the two ultra-runners’ blood glucose concentration. Empirical evidence supports the assumption that a hypoglycemic state was present, and treatment of presumed hypoglycemia without definitive proof is supported by recent ultra-endurance consensus guidelines.7 At this time the theorized pathophysiology of the documented hypothermia is unable to be definitively proven. A rectal temperature, which is considered the gold standard,7 would have provided a more accurate measurement as axillary and oral thermometers may underestimate core temperatures. Rectal thermometer probes (like glucometers) were not available at the time of the patient’s care. Ultramarathon patients are often cared for in resource scarce wilderness settings, where a lack of adequate privacy and absence of multiple thermometers or glucometers per medical team members make empiric diagnoses and subsequent treatment a necessity.

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