Heat-related illness in the African wilderness

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Wilderness heat-related illnesses span a variety of conditions caused by excessive or prolonged heat exposure, and/or the inability to compensate adequately for increased endogenous production during strenuous outdoor activities. Despite management of well-known risk factors, such as lack of fitness or acclimatisation, dehydration, underlying illness and certain medications, even highly trained individuals may exceed their physiological capability to dissipate increased core temperature. Heat illnesses range from benign cramps to the more concerning heat syncope and exercise-associated collapse (with or without hyperthermia), and culminate in life-threatening heat stroke.

The differential diagnosis in the wilderness is broad and should include exercise-associated hyponatraemia with or without encephalopathy. Clinical guidelines for wilderness and hospital management of these conditions are available. Field management and evacuation are based on severity, and include cooling, rehydration and assessment of core temperature and serum sodium, if possible. Hyponatraemia should be corrected with the use of oral or intravenous hypertonic saline, depending on whether the patient can safely take oral fluids. Hospital management may include aggressive and potentially invasive cooling, careful assessment for organ dysfunction, and intensive multi-organ support, if required. Paracetamol, non-steroidal anti-inflammatory drugs and dantrolene should not be used.

Scenario
A fit-looking 23-year-old male participant in a summer mountain trail run collapses 3 km from the end of the 25 km race. Despite the altitude, ambient temperatures are warm, and the participants have been exposed to the full sun for several hours. On your first assessment, he is very confused, with a weak, rapid pulse (160 beats per minute). His skin is wet and cool. A passing member of the same running club alleges that he is not known to have any medical conditions. You notice that his water bottles are both empty. As fellow runners help you carry him to the nearby medical-aid station, a rapidly expanding differential diagnosis fills your thoughts.

Background
Wilderness heat-related illnesses span a continuum of medical problems caused by the generation of, and/or exposure to, excessive heat in the wilderness environment. They can range from minor annoyances to life-threatening conditions. Endogenous heat production is directly dependent on the sum of that from inescapable sources, such as basal cellular metabolism and cardiorespiratory work, and heat generated by physical exertion. All heat exchange (gain or loss) occurs through four mechanisms: conduction, convection, radiation and evaporation. Conduction refers to the direct transfer of heat from one object to another through contact. Convection is heat gain or loss due to movement of a fluid (such as air); a convection oven or wind chill from an Antarctic breeze are similar examples. Radiation describes transfer of heat through electromagnetic waves, epitomised by the heat of the sun on skin. Evaporation is the loss of heat induced by the phase change from liquid to gas, which is dramatically affected by ambient vapour pressure or relative humidity.

Many animals have developed ingenious strategies to adapt their physiology to control heat gain or loss. Indeed, it is hypothesised that particular human traits – our upright stance, complex bony structure of the foot, nuchal ligament, relatively hairless skin and copious capacity for perspiration – evolved to give us an advantage in chasing down prey in the prehistorical African heat.[1] Today, however, humans are unique in their ability to adapt the environment to control their physiology through behaviour and material means: use of clothing, shelter, and devices such as heaters and air conditioners. Despite this, we cannot alter the laws of thermodynamics, and people in the wilderness are subject to the same exposure and thermal exchange mechanisms as our primitive ancestors. Fortunately, the development of modern science, clothing technology, and an understanding of physiology allow us to function successfully in environments of extreme heat under many adverse circumstances (Fig. 1).

Risk factors traditionally considered to increase our susceptibility to the deleterious effects of heat stress include obesity, lack of acclimatisation to hot environments, poor physical fitness, extremes of age, underlying illness (e.g. cardiac conditions, hyperthyroidism), dehydration, certain medications (e.g. beta blockers, anticholinergics,
and diuretics) and illicit drugs (e.g. amphetamines, cocaine, and Ecstasy).

Heat-related illnesses are usually preventable. Mitigating factors include improved physical fitness, heat acclimatisation, selection of appropriate clothing, maintaining adequate hydration, sufficient rest periods, and enhancing evaporative cooling. These may be recalled using the CHAFER mnemonic (Table 1). However, it is important to appreciate that exertional heatstroke and other forms of heat illness can occur in spite of the optimal control of these factors. Similarly, it is normal for well-adapted individuals to experience an increase in core temperature above ‘normal’ values during exercise, even in the setting of low environmental temperatures (Fig. 2). Therefore, core temperature alone is not a reliable indicator of heat illness or injury in the absence of other signs, and illness only occurs when physiological compensation to balance heat gain/production with heat loss fails.

**Definition and spectrum of heat-related illnesses**

Heat-related illnesses are not neatly delineated conditions, but rather occur as a spectrum – from benign to catastrophic. To better understand the pathophysiology of heat illness, it is essential to clarify the definitions thought to encompass these entities.

Classifying the manifestations into discrete categories, however, is useful to ascribe the correct urgency of management and level of severity.

**Heat cramps**

Heat cramps are muscle spasms that typically occur in the large muscle groups of the leg, and may be debilitatingly severe. The precise aetiology is still being debated, but is thought to be a combination of calcium and other electrolyte derangements at tissue level, plasma volume depletion, tissue acidosis in the light of respiratory alkalosis from hyperventilation, and altered neuromuscular control related to exertion. As these cramps can occur without exposure to a warm environment and typically resolve rapidly, they are often considered not to be true heat illness.

**Heat rash and sunburn**

Prickly heat or miliaria rubra is infrequently seen in the South African (SA) context, but occurs in humid or tropical climates owing to exocrine sweat duct blockage during copious or prolonged sweating. It presents as an erythematous pruritic rash with papules, usually on the neck, trunk and limbs, but sparing hands and feet. While heat rash alone is more of an irritation than a serious illness, the reduction in sweat production by duct obstruction can greatly reduce efficiency of evaporative heat loss and predispose the sufferer to more severe injuries. Sunburn – epidermal and dermal damage due to exposure to ultraviolet radiation – also limits sweat production. In

![Fig. 2. Exercise-associated hyperthermia, with routinely raised core body temperatures during periods of physical effort (as shown by raised heart rate) despite frigid conditions during a 7-day mountaineering expedition in Patagonia. (Data – R Hofmeyr.]

| Table 1. Mitigating/preventive factors for heat-related illness – CHAFER |
|--------------------------|-----------------------------------------------------------------|
| **CHAFER**               | **Mitigating/preventive factors**                               |
| **Clothing**             | Clothing should provide skin protection, reflection of solar radiation, and encourage evaporative cooling. Loose-fitting, light-coloured, wicking and/or porous items are ideal. Protection for the head, face and neck should be included. |
| **Hydration**            | Evaporative cooling through sweating is essential to maintain heat loss during exertion in warm environments, and can approach 1 000 mL/h in acclimatised individuals. Prehydration and drinking to thirst are effective measures. Imbibing cold fluids improves comfort and may aid heat loss. |
| **Acclimatisation**      | Gradual exposure to increasing levels of exertion in a hot environment induces heat acclimatisation over a period of days to weeks. The predominant mechanism is an increase in quantity and efficacy of sweat production, improving evaporative cooling. Acclimatisation can be accomplished by 60 - 90 minutes of exercise in the heat per day, ideally over a period of 1 - 2 weeks. Benefits are typically lost within a fortnight of withdrawal from a continuous exposure stimulus. |
| **Fitness**              | Physical fitness allows more efficient effort, reducing the rate of endogenous heat production at the same level of activity compared with a less fit individual. It increases the effectiveness and rate of acclimatisation to hot environments, and reduces body fat, which in turn allows more efficient heat loss. |
| **Evaporative cooling**   | In addition to remaining well hydrated to allow for evaporative cooling through sweating, dipping or soaking clothing in water can greatly increase cooling by this mechanism. |
| **Rest**                 | Rest breaks during acclimatisation, strenuous exertion, and the hottest periods of the day reduce the risk of heat illness. This is particularly effective if rest can be taken in a cooler area. |

CHAFER = clothing, hydration, acclimatisation, fitness, evaporative cooling, rest.
its severe form, it can result in a systemic inflammatory response manifesting with headache, nausea, fever or rigors, and delirium.

**Heat syncope and exercise-associated collapse**

This condition is traditionally defined as a transient collapse or brief loss of consciousness occurring immediately after strenuous exercise in a warm environment. The patient usually rapidly regains clear consciousness shortly after becoming supine, or with raising of the legs. It is often seen in athletes, where it is possibly a manifestation of exercise-associated syncope. It commonly occurs immediately after completion of intense exercise. There are sudden decreases in venous return when skeletal muscle contraction is halted and blood pools in the lower extremities, in the context of exercise-induced peripheral vasodilation and decreased systemic vascular resistance. This may be compounded by an abnormal baroreflex response, Bezold-Jarisch or Barcroft-Edholm reflex. As this condition can occur during intense exercise in the absence of hot conditions, it is frequently termed exercise-induced postural hypotension.

**Heat exhaustion**

While heat exhaustion is understood to be a manifestation of intravascular volume depletion and dehydration caused by prolonged effort in a thermally stressful environment, many in the sports medicine community believe that heat exhaustion is a more severe form of exercise-associated collapse or exercise-induced postural hypotension, and not a separate entity. In the wilderness environment, it is typified by symptoms of fatigue, effort intolerance, weakness, headache, nausea, tachycardia and sometimes dyspnoea. The patient may have a warm or cool skin, with profuse sweating. Core temperature is either normal or mildly elevated (<39°C). Critically, there is no persistent change in mental state. If altered mentation remains for more than a few minutes after the patient is supine, with legs raised, it should be treated as heat stroke.

**Exercise-associated hyperthermia**

Raised core body temperature (<40°C) that results from strenuous exertion or exercise may be a normal consequence of increased endogenous heat production, and can occur in cold environments, particularly in athletes exercising at or close to maximum capacity (Fig. 1). Where exercise-associated hyperthermia leads to collapse or altered level of consciousness, it indicates the development of heat exhaustion or heat stroke. In athletes, this typically occurs over medium distances (5 - 21 km). Previous occurrence is a risk factor.

**Heat stroke**

Heat stroke is a true medical emergency. The defining characteristics are a core temperature of >40°C and encephalopathy, manifested by altered mental status. While often distinguished as occurring as a result of prolonged exposure to high ambient temperatures (classic heat stroke) or pathological hyperthermia during exercise (exertional heat stroke), it is most commonly a combination of the two in the context of wilderness medicine. The patient may be sweating, but often has a dry skin. Clinical signs include confusion, disorientation, decreased level of consciousness, inappropriate behaviour, ataxia or collapse, tachypnoea, and tachycardia, or the patient may be in cardiovascular shock on presentation. Dysrhythmias can occur. Risk of brain, renal, hepatic and other organ damage depends on duration of hyperthermia, with systemic protein denaturation beginning at >42°C. The patient may develop a systemic inflammatory response syndrome (SIRS) owing to cytokine and heat-shock protein release, which mimics sepsis. Hypoperfusion of the gastrointestinal tract is exacerbated by inflammatory-mediated increase in mucosal and vascular permeability, leading to endotoxin migration from the gut to the circulation. This is thought to lead to the multi-organ failure that causes death in severe or untreated cases.

**Exercise-associated hyponatraemia**

While not specifically a heat-related illness, exercise-associated hyponatraemia (EAH) often occurs in the setting of prolonged effort in hot conditions. It is therefore important to understand the condition, and to consider it as a differential diagnosis. Defined as a serum sodium concentration <135 mmol/L within 24 hours of prolonged physical activity, it is believed to be caused by an interplay between excessive fluid intake (especially hypotonic fluids such as clear water) and reduced water excretion under the influence of antidiuretic hormone due to the physical stress imposed by exercise. Of these factors, overhydration (>1,500 mL/h) is considered the most prominent cause. While most cases are asymptomatic and go undetected, EAH can cause fatigue, nausea/vomiting, headache, dizziness, blurred vision and tachycardia. Severe cases can result in cerebral oedema (even non-cardiogenic pulmonary oedema) and confusion, decreased level of consciousness, and seizures. In these cases, it is termed exercise-associated hyponatraemic encephalopathy (EAHE).

**Wilderness medicine guidelines for heat-related illnesses**

The Wilderness Medical Society (WMS) has published practice guidelines for wilderness emergency care, heat-related illnesses, and EAH, most recently updated in 2014. These peer-reviewed, expert-consensus guidelines form an excellent foundation upon which local recommendations can be made for the African context, drawing from local experts in fields such as wilderness, emergency and sports medicine.

**Clinical diagnosis and temperature measurement**

The practitioner is usually alerted to the initial presence of heat-related illness by patient symptoms. In the wilderness, a presumptive diagnosis is made on the clinical signs described above, in the presence of heat exposure and/or excessive endogenous production. However, as hyperthermia and heat stroke can exist in the absence of excessive ambient temperatures – even in patients with cool extremities and skin – contemporaneous measurement of core body temperature is very valuable in excluding or confirming the diagnosis, assessing severity, and monitoring response to treatment. Unfortunately, accurate core temperature measurement in the field is challenging. Invasive gold-standard measurements (pulmonary artery catheter or oesophageal thermistor) are infrequently available, and skin, oral or infrared tympanic measurements lack accuracy. A pragmatic approach is to treat lucid patients according to their perceived temperature (as they do not have heat stroke if fully conscious and orientated), use a rectal thermometer for patients with altered consciousness, and insert an oesophageal thermistor, when available, in unconscious patients who have an advanced airway in place.
Field management

Severity of outcome of heat-related illness is directly proportionally related to duration of exposure. Treatment should therefore be initiated in the field, both to limit the progression of severity and to limit duration. Fortunately, there are many interventions that can be commenced in resource-constrained settings that will improve patient outcome (Table 2).

Field testing of serum sodium is available using compact, battery-powered point-of-care devices, such as the i-STAT analyser (Abbott Point of Care, USA). However, in very hot environments, these devices often have a limited operation temperature range (e.g. 16 - 30°C for the i-STAT), and the device may have to be kept cool in the field to guarantee function when required.

Hospital management

Cases of suspected heat-related illness should always be managed urgently on admission to hospital. In addition to swift primary assessment and any related advanced life-support interventions, a reliable (rectal or oesophageal) core temperature and serum sodium measurement should be obtained as a matter of priority. Intravenous access should be established. It is valuable to obtain haematological, electrolyte, creatinine kinase, renal and liver function tests at the same time. Patients with altered mental status and a core temperature of >39°C must be urgently cooled, using methods described below. Invasive cooling using bladder, gastric, pleural or peritoneal lavage can be considered if the patient does not respond rapidly. Seizures should be controlled with a benzodiazepine or propofol, which may necessitate intubation and ventilation. Treat hypotension or shock with fluid resuscitation, but avoid hypotonic fluids, especially if the serum sodium level is unknown. If EAHE is diagnosed, up to three 100 mL boluses of 3% hypertonic saline at 10-minute intervals are advocated. There is no risk of central pontine myelinolysis with this therapy in the context of EAHE.

Serum potassium and calcium should be assessed and corrected if abnormal. If there is biochemical evidence of rhabdomyolysis, promoting positive fluid balance, diuresis with furosemide, and alkalisation of urine with sodium bicarbonate can be considered. In contrast to fever, there is no role for paracetamol and non-steroidal anti-inflammatory drug use in heat stroke, which may cause further harm. Dantrolene has also not been shown to contribute to management. Admission to an intensive care unit (ICU) with renal dialysis and management for multi-organ failure may be required. The treating clinicians should be alert for acute respiratory distress syndrome, liver or kidney injury, coagulopathy and gastrointestinal ischaemia.

Scenario resolution

You consider the potentially differential diagnosis, including presentation of underlying febrile illnesses such as malaria, hypoglycaemia, drug toxicity or intoxication, cardiac ischaemic events, epilepsy, an occult head injury or EAHE. However, the hot conditions and strenuous exertion make heat exhaustion or heat stroke very likely.

Table 2. Field management of heat-related illness

<table>
<thead>
<tr>
<th>General interventions</th>
<th>In all suspected heat-related illness, decreasing physical effort or enforcing rest, moving the patient to a cool environment (or taking steps to shade the patient and encourage evaporative cooling), encouraging oral hydration if thirsty, and considering replacement of electrolytes are essential steps to enhance recovery.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat cramps</td>
<td>Rest and stretch the involved limb and muscle groups. Encourage hydration with electrolytes. Consider calcium/magnesium supplementation.</td>
</tr>
<tr>
<td>Heat rash</td>
<td>Promote a cool environment and improved personal hygiene. Application of lotions will aggravate the condition, although some advocate a topical corticosteroid in severe cases. Acclimatisation and avoidance of sweating will reduce further incidences.</td>
</tr>
<tr>
<td>Sunburn</td>
<td>Prevention of further exposure using barriers (clothing and regularly applied high-factor sunscreen) is essential. Hydrating lots and oral anti-inflammatory may reduce severity.</td>
</tr>
<tr>
<td>Heat syncope</td>
<td>Cool the patient where possible; place supine and elevate legs. Once recovered, decrease effort and encourage frequent rest. Avoid aggressive intravenous and other invasive therapies.</td>
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<tr>
<td>Heat exhaustion</td>
<td>Cool environment, rest, rehydration (typically oral), and consider electrolytes (oral). Assess core temperature early if there is any evidence of change in mental status. Evacuation for severe cases or those who respond poorly.</td>
</tr>
<tr>
<td>Symptomatic wilderness hyperthermia/heat stroke</td>
<td>Provide advanced life support as indicated. Administer oxygen if available and saturations are &lt;92%. Aggressive cooling using any available means should be initiated, including shading the patient from sunlight, removing excess clothing, soaking, dousing or partially submerging the patient in cold water, fanning, cold IV fluids, or even an ice bath. Covering the patient with a soaked blanket/sheet with continuous fanning is an effective method, using convective and evaporative cooling. Ice packs can be applied to the neck, axilla and groin, but their use and effectiveness have been challenged. No paracetamol or NSAIDs should be administered. Invasive/rectal temperature monitoring should be established. Cool until 39°C and then observe; beware of rebound hyperthermia, but also excessive cooling. If using cold water or ice baths to cool the patient, it is effective to continue until the conscious patient begins to complain of cold. All patients should be evacuated to definitive medical care urgently.</td>
</tr>
<tr>
<td>EAH and EAHE</td>
<td>EAH/EAHE should form part of the differential diagnosis for heat-related illnesses in the field. Ideally, field testing of serum should be used, as this guides the selection of fluid (hypo-, iso- or hypertonic) used for treatment. In the absence of measurement, hypotonic or isotonic IV fluids should be avoided unless there is a clear indication (e.g. hypotension/shock). If hyponatraemia (serum sodium &lt;135 mmol/L) can be demonstrated in the field, oral or IV hypertonic saline (100 mL 3% NaCl) can be administered every 10 minutes until 3 doses have been completed or neurological symptoms improve. An oral mixture of hypertonic saline can be created by adding 3 - 4 stock cubes to half a cup of water.</td>
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</table>

EAH = exercise-associated hyponatraemia; EAHE = exercise-associated hyponatraemic encephalopathy; IV = intravenous; NSAIDs = non-steroidal anti-inflammatory drugs.
At the medical station, you elevate the athlete’s legs, rapidly commence cooling by soaking his clothing in cool water and encouraging evaporation by fanning, and obtain a rectal temperature of 41.2°C. Recognising the medical emergency, you establish IV access, and recognising the risk of hyponatraemia, start isotonic fluid resuscitation. You request an urgent evacuation, even though the patient is gradually becoming more lucid. On arrival at the emergency unit, he is found to have significant rhabdomyolysis and dehydration in addition to heat stroke, but despite spending a night in the ICU, he avoids more serious consequences because of your swift interventions.