Heat Illness. A Report of 45 Cases from Hong Kong

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SUMMARY: During the summer of 1985, 45 soldiers with heat illness were admitted to the British Military Hospital, Hong Kong. Twelve had severe heat stroke, the remainder heat exhaustion. This paper discusses the management and prevention of heat illness in the military context.

Introduction
The health hazard posed by high environmental temperature and humidity is frequently underestimated, yet, in the USA alone over 4000 heat related deaths are recorded annually 1. Although the burden of mortality and morbidity falls on the elderly, young fit people, particularly soldiers undergoing vigorous training, may suffer severely 2. Heat illness has a broad spectrum of severity. Mild cases (heat exhaustion) recover quickly with cooling and rehydration, whilst those with core temperatures greater than 41°C or cardiovascular or neurological complications (heat stroke) suffer a severe multisystem disorder with a mortality exceeding 50% 2-9.

Heat illness is caused by failure of the normal heat dissipation mechanisms allowing an excessive rise in core temperature. Once the core temperature exceeds 41°C protein denaturation begins 7. Failure of adequate heat transfer to the environment is usually due to high environmental temperature or humidity although other factors such as excessive clothing and lack of air movement are important 9. Other factors predisposing to heat illness are vigorous exercise, age, poor acclimatization, alcohol abuse, obesity, febrile illness, heart disease, drugs (especially atropine and phenothiazines) and inadequate fluid intake 2-12.

The hot humid conditions which prevail during the Hong Kong summer have long been recognised as posing an environmental heat threat to military operations. The British Army uses the Wet Bulb-Globe Temperature Index (WBGT Index) to predict the severity of the heat threat 13. It is calculated by the addition of 70% of the wet bulb temperature to 20% of the black bulb temperature plus 10% of the dry bulb temperature. The implications of the WBGT Index for military operations in Hong Kong are shown in Table 1.

This paper reviews all the heat illness casualties treated in the British Military Hospital (BMH) Hong Kong during 1985.

Method
All heat casualties admitted to BMH Hong Kong during 1985 were studied prospectively. They were assessed on admission according to a detailed protocol and assigned to one of two groups:

**a. Heat Exhaustion**
Defined as a rectal temperature of under 41°C, normal peripheral perfusion, normal blood pressure and clear mentation or mild confusion. Immediate investigations included thick film examination for malarial parasites, serum urea and electrolytes, white cell count and packed cell volume.

**b. Heat Stroke**
Defined as a rectal temperature of over 41°C or lesser degrees of pyrexia in association with severe confusion, unconsciousness, hypotension or peripheral circulatory failure. Immediate investigations included thick film examination for malarial parasites, serum urea and electrolytes, blood glucose, prothrombin time, fibrinogen degradation products, liver function tests, packed cell volume, white cell count, platelet count, serum creatine phosphokinase and hydroxybutyrate dehydrogenase, a chest X-ray and a 12 lead electrocardiogram. Further investigations such as blood gas analysis were done as indicated. Treatment was standardised using a predetermined heat illness protocol.

*Also Comd Adv General Practice Hong Kong & SMO Gurkha Fd Force
Table I

<table>
<thead>
<tr>
<th>Heat Index</th>
<th>Acclimatised troops</th>
<th>Unacclimatised troops</th>
</tr>
</thead>
<tbody>
<tr>
<td>26°C (Green)</td>
<td>Normal Work</td>
<td>Use discretion</td>
</tr>
<tr>
<td>28°C (Amber)</td>
<td>Use discretion</td>
<td>Avoid work for the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>first 7 days</td>
</tr>
<tr>
<td>30°C (Red)</td>
<td>Use discretion</td>
<td>Avoid work for the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>first 21 days</td>
</tr>
<tr>
<td>32°C or above</td>
<td>Avoid heavy exercise</td>
<td>Avoid heavy exercise</td>
</tr>
<tr>
<td>(Black)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

On recovery all British and Gurkha soldiers were questioned about water discipline and their general knowledge of the environmental heat threat.

Results
A total of 45 male soldiers with heat illness were admitted to the study. All the casualties occurred during the hot humid summer between August and early November. They had all collapsed during or immediately after a period of vigorous physical exercise and all but two were evacuated to hospital by helicopter. They fell into three well defined groups which will be considered separately.

Group I: Casualties from an Australian Army Exercise
During August (the hottest and most humid month in Hong Kong) the Australian Army held a company strength exercise involving an assault on a hill. The exercise began in the late morning when the WBGT Index was AMBER. Twenty-two men collapsed during the assault. They were doused with water, but not stripped, then evacuated to hospital by helicopter. All 22 arrived within a two hour period. None had their body temperature measured in the field. On arrival they were all lucid or only slightly confused, normotensive and normothermic. All were given a litre of normal saline intravenously and allowed free access to iced Dioralyte (Armour Pharmaceuticals: one sachet made up to 200 mls with water contains sodium chloride 200 mg, potassium chloride 300 mg, sodium bicarbonate 300 mg and glucose 8 g.). They all recovered quickly and were discharged within 48 hours.

Group II: Casualties from Exercise Bulldog Patrol
Between the seventh and tenth of October Gurkha Field Force held a major exercise in the New Territories. The exercise finished with a forced march followed by a 2½ mile speed march carrying full equipment. The WBGT Index for the first two days of the exercise was unavailable due to a machine fault, Amber on Day 3 and Green on Day 4. A total of 18 serious heat casualties occurred during or immediately after the speed march. All were taken to a nearby medical centre where they were stripped, covered in a wet sheet and vigorously fanned. Intravenous rehydration was started immediately using Hartmann's solution. The casualties were then rapidly evacuated to hospital by helicopter. The mean delay between arrival at the medical centre and admission to hospital was 45 minutes (range 15-60 minutes).

Clinical Features (See Table II)
All had a short prodrome of headaches, cramps, nausea or vomiting, dizziness or irrational behaviour before collapsing. Eleven were seriously ill with established heat stroke. Of these, three were very confused, one was delirious and two were unconscious and unresponsive to painful stimuli. Three had rectal temperatures of 42°C or higher. First aid at the point of collapse and the cooling effect of the helicopter flight produced a fall in rectal temperature in every case.

There was a striking difference in the clinical condition of the British soldiers compared with the Gurkhas despite similar thermal insult. All the British soldiers were, on arrival, mentally clear, normotensive and well perfused peripherally. Of the Gurkhas by contrast six had severely impaired mental function (two in coma) and although normotensive or near normotensive six were grossly vasoconstricted with cold cyanosed peripheries despite an elevated core temperature.

Patients with rectal temperatures above 38°C were cooled by tepid sponging and vigorous fanning. Once the rectal temperature fell below 38°C active cooling was stopped to prevent an afterdrop in core temperature. Rehydration was with intravenous normal saline (infused cooled for patients still pyrexial) followed as soon as possible by oral Dioralyte. All were significantly dehydrated (See Table III) and some did not produce urine until more than two litres of intravenous fluid had been infused. In patients with depressed consciousness or peripheral circulatory failure urinary catheters were
There were no deaths.

Hartmann’s solution before arrival at hospital.

Biochemical Data

To more invasive techniques such as cold peritoneal rehydration and all were fully recovered and discharged within three days. The Gurkhas by contrast had an peripheral circulatory failure which could not be reversed by rehydration. Incremental doses of chlorpromazine given as for shiver suppression produced sufficient cutaneous vasodilatation to allow cooling without having to resort to more invasive techniques such as cold peritoneal dialysis.

The British soldiers responded quickly to cooling and rehydration and all were fully recovered and discharged within seven days (range 2-11 days). There were no deaths.

Biochemical Data (See Table III)

All patients had received at least one litre of Hartmann’s solution before arrival at hospital. Serum sodium and potassium were normal. Serum urea was modestly raised and normalised in every case after rehydration. The mean serum urea in the Gurkhas was slightly but significantly higher than in the British soldiers (mean urea of 8.1 mmol/l versus 6.8 mmol/l. T=0.8 p<0.05). Liver function tests and muscle enzymes were measured in men with heat stroke. Serum bilirubin was normal in all cases. Aspartate transaminase (normal 5-50 iu/l) was raised in 10 (maximum value 108 iu/l), alanine transaminase (normal 5-50 iu/l) was raised in one to 85 iu/l, hydroxybutyrate dehydrogenase (normal 52-222 iu/l) was raised in seven (maximum value 307 iu/l) and creatine phosphokinase (normal 45-390 iu/l) was raised in three (maximum value 673 iu/l).

Haematological Data (See Table III)

All had modest neutrophil leucocytosis; the mean white cell count was 14.0 x 10⁹/l (range 6.8- 25.2 x 10⁹/l). None had a clinically important bleeding diathesis although three had prolongation of prothrombin time (up to 35 seconds) and one had slightly elevated fibrinogen degradation products (up to 40). These abnormalities disappeared within 48 hours. Platelets remained normal in every case.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Diagnosis Race</th>
<th>Temperature °C Medical Centre</th>
<th>Temperature °C BMH</th>
<th>Mental State</th>
<th>Peripheral Vasoconstriction</th>
<th>Peripheral Cynosis</th>
<th>BP at BMH (mm Hg)</th>
<th>ECG</th>
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<tr>
<td>23</td>
<td>HS G</td>
<td>40.5</td>
<td>39.0</td>
<td>Coma</td>
<td>Severe</td>
<td>Severe</td>
<td>120/70</td>
<td>Normal</td>
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<tr>
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<td>38.0</td>
<td>Normal</td>
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<td>Absent</td>
<td>110/70</td>
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<tr>
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<td>HS G</td>
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<td>39.0</td>
<td>Confused</td>
<td>Absent</td>
<td>Absent</td>
<td>80/50</td>
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</tr>
<tr>
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<td>42.4</td>
<td>37.5</td>
<td>Normal</td>
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<td>Severe</td>
<td>110/70</td>
<td>Abnormal</td>
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<td>40.6</td>
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<td>Severe</td>
<td>100/60</td>
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<td>Severe</td>
<td>Severe</td>
<td>125/70</td>
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<tr>
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<td>Absent</td>
<td>100/60</td>
<td>Normal</td>
</tr>
<tr>
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<td>37.5</td>
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<td>Severe</td>
<td>Severe</td>
<td>120/90</td>
<td>Normal</td>
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<tr>
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<td>Absent</td>
<td>130/70</td>
<td>Normal</td>
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<td>Absent</td>
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<td>Normal</td>
</tr>
<tr>
<td>33</td>
<td>HS G</td>
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<td>37.0</td>
<td>Confused</td>
<td>Severe</td>
<td>Severe</td>
<td>130/80</td>
<td>Normal</td>
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<tr>
<td>34</td>
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<td>37.5</td>
<td>Normal</td>
<td>Absent</td>
<td>Absent</td>
<td>130/70</td>
<td>Normal</td>
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<tr>
<td>35</td>
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<td>37</td>
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<td>Absent</td>
<td>120/70</td>
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<tr>
<td>38</td>
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<td>Normal</td>
<td>Absent</td>
<td>Absent</td>
<td>115/80</td>
<td>Normal</td>
</tr>
<tr>
<td>39</td>
<td>HE G</td>
<td>39.5</td>
<td>38.5</td>
<td>Normal</td>
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<td>HE G</td>
<td>38.5</td>
<td>37.5</td>
<td>Normal</td>
<td>Absent</td>
<td>Absent</td>
<td>110/80</td>
<td>Normal</td>
</tr>
</tbody>
</table>

HS - Heat Stroke  G - Gurkha
HE - Heat Exhaustion  B - British
At last, an exclusively topical antibiotic.

Bactroban
mupirocin
for acute primary skin infections.

**PRESCRIBING INFORMATION**

**USES. Action:** Bactroban is a topical antibacterial agent, active against those organisms responsible for the majority of skin infections, e.g., Staphylococcus aureus, including methicillin-resistant strains, other staphylococci, streptococci. It is also active against Gram-negative organisms such as Escherichia coli and Haemophilus influenzae.

**Indications:** Acute primary bacterial skin infections, e.g., impetigo and folliculitis.

**DIRECTIONS FOR USE:** Adults and children: Apply up to three times a day, for up to 10 days. (There is no long-term experience of Bactroban in humans).

**CONTRA-INDICATIONS, WARNINGS ETC:** Contra-indications: Hypersensitivity to Bactroban or otherointments containing polyethylene glycols. This Bactroban ointment formulation is not suitable for ophthalmic or intra-nasal use. Use in pregnancy: There is inadequate evidence of safety to recommend the use of Bactroban during pregnancy. Precautions: Avoid contact of Bactroban Ointment with the eyes. Use Bactroban Ointment with caution in patients with moderate or severe renal impairment.

**SIDE-EFFECTS:** Some minor, localised side-effects, such as burning, stinging and itching have been reported.

**PRESENTATION AND BASIC NHS PRICE** (Correct at February 1986): Bactroban Ointment: mupirocin 2% w/w. £3.45 per 15g tube in a sealed carton.

Further information on Bactroban, a product of British research, is available on request to Beecham Research Laboratories, Brentford, Middlesex, TW8 9BD.
Table III
Biochemical and haematological data from Bulldog Patrol heat casualties

<table>
<thead>
<tr>
<th>Patient</th>
<th>Serum Urea mmol/L</th>
<th>Serum Sodium mmol/L</th>
<th>Serum Potassium mmol/L</th>
<th>Packed cell volume</th>
<th>White cell count x 10^9/L</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>23</td>
<td>24</td>
<td>25</td>
<td>26</td>
<td>27</td>
</tr>
<tr>
<td>Urea</td>
<td>5.8</td>
<td>4.7</td>
<td>7.2</td>
<td>9.4</td>
<td>6.7</td>
</tr>
<tr>
<td>Sodium</td>
<td>144</td>
<td>139</td>
<td>146</td>
<td>138</td>
<td>139</td>
</tr>
<tr>
<td>Potassium</td>
<td>4.7</td>
<td>5.0</td>
<td>4.5</td>
<td>4.6</td>
<td>5.3</td>
</tr>
<tr>
<td>Packed cell volume</td>
<td>43%</td>
<td>46%</td>
<td>48%</td>
<td>46%</td>
<td>43%</td>
</tr>
<tr>
<td>White cell count x 10^9/L</td>
<td>9.5</td>
<td>17.2</td>
<td>11.1</td>
<td>12.1</td>
<td>11.0</td>
</tr>
</tbody>
</table>

Normal Values
Urea 2.8 - 7.2 mmol/L  
Sodium 130 - 142 mmol/L  
Potassium 3.5 - 5.00 mmol/L  
Packed cell volume 47 ± 7%  
White cell count 4 - 11 x 10^9/L
Electrocardiography (See Table II)

The ECG remained normal in 12. Six patients (all Gurkhas) had evolving ECG changes which were maximal between 48-72 hours. These changes comprised inferior ST segment depression and T wave inversion or anterior ST segment elevation with upward coving and T wave flattening or inversion in a manner reminiscent of acute subendocardial infarction. The changes were not associated with chest pain or arrhythmias and had normalised in all but one case by 14 days. In all six, resting myocardial thallium scans were carried out during the third week but no abnormalities were detected.

Water Discipline

The British soldiers had all drunk between two and four litres of fluid during the forced march and had had compulsory stops for cooling and fluid replenishment. The Gurkhas had all drunk before the march and had not included water stops during the march and had been advised against drinking during heavy exercise. Seven had drunk no fluid at all while four had drunk up to \( \frac{1}{2} \) litre.

Knowledge of heat illness

The British soldiers were all aware of the heat hazard but underestimated its magnitude. They had all heard of the heat illness immediate action drill, although in four of the seven their knowledge was inadequate. The Gurkhas seemed to be only dimly aware of the environment heat hazard and none knew the heat illness immediate action drill.

Pre-existing Illness

Three of the heat victims had “flu-like” symptoms within 24 hours of the exercise but none had reported sick. One Gurkha (case 23) suffered a relapse of fever on the day after admission. Examination revealed a small eschar and splenomegaly. A clinical diagnosis of scrub typhus was made and he responded rapidly to doxycycline.

Disposal

Following recovery all were subject to restrictions determined by the severity of their illness:

- Heat exhaustion: no strenuous exercise for two weeks
- Heat stroke: no strenuous exercise for four weeks
- Abnormal ECG: medically downgraded to P3 for three months with a prohibition on strenuous exercise during that period.

Group III. Sporadic Cases (See Table IV)

Five men not involved in major exercises suffered heat illness. All had been exercising vigorously before collapsing. Four had heat exhaustion and recovered quickly on cooling and rehydration; they had taken large volumes of fluid while exercising. One Gurkha (case 45) had severe heat stroke; he collapsed after a run during which he had drunk nothing. He was not given first aid at the point of collapse and did not reach the medical centre until half an hour later. He was then deeply unconscious with a rectal temperature of 38.5°C. He was stripped, cooled and given a litre of intravenous Hartmann's solution before being taken by helicopter to hospital. Despite rapid cooling and rehydration in

<table>
<thead>
<tr>
<th>Patient</th>
<th>Diagnosis</th>
<th>Serum Urea mmol/l</th>
<th>Serum Sodium mmol/l</th>
<th>Serum Potassium mmol/l</th>
<th>Packed cell volume</th>
<th>White cell count ( \times 10^9/l )</th>
<th>Race</th>
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<tr>
<td>41</td>
<td>HE</td>
<td>6.1</td>
<td>138</td>
<td>4.2</td>
<td>39</td>
<td>18.3</td>
<td>B</td>
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<tr>
<td>42</td>
<td>HE</td>
<td>8.5</td>
<td>144</td>
<td>4.7</td>
<td>55</td>
<td>12.6</td>
<td>B</td>
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<tr>
<td>43</td>
<td>HE</td>
<td>3.8</td>
<td>140</td>
<td>3.8</td>
<td>43</td>
<td>6.8</td>
<td>G</td>
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<tr>
<td>44</td>
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<td>10.6</td>
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<tr>
<td>45</td>
<td>HS</td>
<td>7.8</td>
<td>141</td>
<td>4.1</td>
<td>36</td>
<td>10.3</td>
<td>G</td>
</tr>
</tbody>
</table>

Normal Values

- Sodium 130 – 142 mmol/L
- Urea 2.8 – 7.2 mmol/L
- Potassium 3.5 – 5.0 mmol/L

\( HS = \) Heat Stroke
\( HE = \) Heat Exhaustion
\( G = \) Gurkha
\( B = \) British
Further examination revealed gross truncal ataxia and hospital he remained very drowsy for three days. Further examination revealed gross truncal ataxia and dysarthria. A cerebral CAT scan was normal and viral titres (including Japanese encephalitis) were negative. His CSF was normal except for a slightly elevated protein (1.2 g/l). He gradually recovered over six weeks.

Discussion

The heavy toll of heat casualties provided not only experience in the management of heat illness of all grades of severity but highlighted several problem areas.

1. The Multiple Casualty Situation

Heat illness is not often thought of as a cause of multiple casualties yet its potential in this regard was amply shown by both the Australian and British Army exercises. During the Australian exercise all 22 casualties were admitted within a short time. Their early assessment and management was greatly eased by the existence of a standardised policy (Heat Illness Protocol) for the triage, investigation and management of heat victims. Using the protocol, junior doctors, inexperienced in heat illness, and senior nursing staff could begin essential investigations and treatment without having to wait for advice from a senior colleague.

The multiple casualty situation during Bulldog Patrol was different and more complex in that 3-5 seriously ill patients arrived together on each of four consecutive days. A senior doctor was assigned to the intensive care unit to supervise resuscitation. As the intensive care facility became full, further direct admissions were stopped. New arrivals were assessed in an adjacent room by another experienced doctor before being transferred to the general ward or intensive care for resuscitation and treatment. Here again the standardised Heat Illness Protocol was invaluable in providing guidance for junior doctors and nurses at a time when senior colleagues were fully occupied with gravely ill patients.

2. Management of Heat Exhaustion in Hospital

The duration of hyperpyrexia is a crucial prognostic factor. Prompt first aid in the field can do much to prevent mortality and permanent neurological sequelae in heat victims. The speed of cooling is more important than the method. During the Australian exercise, sensible measures at the point of collapse probably prevented heat stroke in some men, whilst during Bulldog Patrol more sophisticated measures taken at the nearby medical centre probably saved lives of some of the worst cases. At the most basic level, the victim should be removed to a shaded, cool place, stripped off and fanned. If water is available, dousing the victim enhances cooling. Those capable of drinking should be encouraged to do so, although depressed consciousness and nausea are limiting factors. At a more sophisticated level rapid intravenous rehydration will mitigate the worst effects of heat illness. Transportation to hospital by helicopter has the advantage of speed and produces rapid cooling due to draught. It was our impression that the more open Scout helicopter produced better cooling than the Wessex. In the absence of a helicopter, a vehicle with a covered roof and open sides will produce effective cooling, particularly if the victim is wrapped in a wet sheet.

3. Management of Heat Exhaustion in Hospital

Providing that victims are rapidly cooled and rehydrated full recovery within 48 hours can be expected. We found stripping, tepid sponging and vigorous fanning to be an effective method of cooling. Although many patients will achieve water and electrolyte homeostasis if given free access to oral fluids (we used Dioralyte) some are intolerant to oral rehydration because of nausea and vomiting. This is especially troublesome in patients who have experienced severe salt depletion. To avoid this we routinely used intravenous rehydration in addition to free oral fluids during the first 12 hours.

4. Management of Heat Stroke in Hospital

All patients with established heat stroke were admitted to the intensive care unit because of the risk of sudden death and the everpresent threat of complex multisystem failure.

Cooling

Stripping, tepid sponging and fanning were used as a routine and were effective in patients without peripheral circulatory failure. Large volumes of chilled normal saline were held in the intensive care unit for early resuscitation of hyperpyrexial patients. Even in the presence of a raised core temperature active cooling invariably produced a highly undesirable heatproducing shiver response. Shivering was abolished by intravenous chlorpromazine 25 mg repeated up to 75 mg. It is vital to delay using chlorpromazine until intravenous fluid repletion is underway to avoid hypotension. Blood pressure must be monitored very frequently.

Recently it has been shown that very rapid cooling can be achieved by placing the victim on an open string mattress and spraying atomised water driven by warm air from above and below. Methods which rely upon surface cooling can not be used alone in hyperpyrexial patients with profound cutaneous vasoconstriction due to peripheral circulatory failure. We found using chlorpromazine as for shiver suppression produced sufficient cutaneous vasodilatation to allow surface cooling so avoiding more dangerous techniques such as cold peritoneal dialysis. Active cooling was stopped once the rectal temperature fell below 38° to avoid an undesirable after drop in core temperature.
Rehydration

All patients were rehydrated with intravenous normal saline. Urinary catheters were passed in patients with depressed consciousness or peripheral circulatory failure to monitor hourly urine flow. Some patients required more than two litres of intravenous fluid before any urine appeared. In our patients central venous pressure monitoring was not necessary although it was available.

Management of Complications

1. Neurological

Confusion is common\(^2\). Three patients in this study were very confused but all recovered without sequelae. One patient was severely disturbed and required sedation for 24 hours with intravenous chlorpromazine. On recovery he suffered troublesome headache for several days. Three patients were unconscious on admission. Coma can develop with remarkable speed in severe heat illness and is an ominous prognostic sign\(^2,3\). All survived although one suffered severe cerebellar dysfunction which persisted for several weeks. Permanent neurological damage may occur and was probably averted in this series by early aggressive treatment\(^2,9,11,17\). In patients with severely disturbed consciousness alternative diagnoses such as cerebral malaria, meningitis, encephalitis or hypoglycaemia should not be forgotten.

2. Cardiovascular

Six patients had profound peripheral circulatory failure with cold cyanosed extremities. Despite normal or near normal blood pressure peripheral circulatory failure in the context of heat stroke implies a grossly reduced cardiac output (under 3 litres/min/m\(^2\)) and is a bad prognostic sign\(^2,18,19\). Its mechanism is complex. With core temperatures above 41°C direct myocardial depression occurs\(^20\) but a more important mechanism appears to be hypovolaemia due to excessive sweating, leakage of fluid across capillary membranes rendered incompetent by heat injury and inadequate fluid intake\(^9,21\). They responded poorly to volume replacement and oxygen therapy. Small repeated doses of intravenous chlorpromazine reduced the intensity of the vasoconstriction and improved urine flow. Specific inotropes such as dopamine were not necessary.

Serial ECG monitoring revealed evolving changes suggestive of subendocardial infarction in six Gurkhas. None had been significantly hypotensive but three had severe peripheral circulatory failure on admission.

This type of electrocardiographic pattern has been described by others and the changes may persist for months\(^3,8,18,22,23\). In fatal cases spotty myocardial necrosis is found\(^3,21,24\) and this type of pathology may have been responsible for the ECG changes in our series. The normal myocardial thallium scans excluded large areas of necrosis but the method is probably too insensitive to detect very small areas of necrosis.

Collateral evidence of infarction can not be obtained from assay of cardiac enzymes as these may be elevated after heavy exertion and in patients with heat stroke without other evidence of myocardial injury\(^25\).

Haematological Abnormalities

A neutrophil leucocytosis is a feature of heat illness and was present in all patients in this series. It has no significance beyond realising that it does not imply bacterial infection\(^11\). Minor derangements of clotting are common but as in this series, they are usually of little clinical importance\(^3,4,8,9,26\). Major bleeding due to disseminated intravascular coagulation has been described and appears to respond to clotting factor replacement and heparin\(^2,6,26\).

Metabolic Complications

Mild metabolic acidosis was common but responded to intravenous fluid replacement, and specific therapy with bicarbonate was not required. Although the fluid and electrolyte status of patients depends upon the quantity of fluids drunk most patients have a combined water, sodium and potassium deficiency\(^2,8\). We routinely used normal saline to replace circulating volume and added potassium chloride to the intravenous fluids as soon as patients began to pass urine. None of our patients developed oliguric or polyuric renal failure although both are well known complications of heat stroke\(^20,27,28\).

The liver is very sensitive to heat injury and biochemical evidence of liver damage is common\(^2,21\). As in our series it is usually of little clinical significance although occasionally it may be severe enough to contribute to death\(^21\).

5. Prevention of Heat Illness

Although it may be impossible to eliminate heat illness from military operations in the tropics much can be done to reduce its prevalence and severity.

WBGT Index. This provides commanders with a rapid assessment of the environment heat hazard (Table 1). It is vital to realise however that indices below BLACK do not equate with safety. Experience in Hong Kong during 1985 has shown an AMBER reading to be associated with a grave hazard.

Acclimatization. Although this was not a factor in our patients it is a well recognised variable in the genesis of heat illness. The mechanism is complex but excessive sweat loss in unacclimatized people undergoing hard physical training may be important\(^9,11\).

Timing of Exercise. Where possible strenuous exercise should be avoided during the hottest period of the day\(^11\). This may not always be possible in the military context but the penalty can be high as may be seen from the heat casualties from the Australian Army Exercise.

Avoiding Dehydration. Heavy exercise in hot weather causes enormous loss of water, sodium and potassium in sweat. Peak sweating rates may exceed two litres per
hour². During strenuous exercise adequate stops for cooling and rehydration are essential if heat illness is to be avoided²,⁹,¹¹,¹². Lack of proper water stops probably explains the extremely severe heat illness suffered by the Gurkha soldiers. Although giving additional salt had been advocated during heavy exercise, it appears that this is not only unnecessary, but potentially dangerous, as it may induce hypokalaemia²,¹¹.

Increasing Knowledge of Heat Illness. The results of the questionnaire filled in by the heat victims from Bulldog Patrol showed a low level of knowledge regarding heat illness particularly with the Gurkhas. Caution however must be used in extrapolating this type of data to the battalions in general. Clearly it is essential that all soldiers are familiar with the heat illness immediate action drill. Commanders at all levels need to be aware of the prodromal symptoms of heat illness such as faints, dizziness, cramps, vomiting or unusual behaviour which although readily reversible with cooling and rehydration, can, if ignored, rapidly progress to fatal heat stroke¹¹. If possible light open clothing should be worn although in the military context this may not always be possible¹.

Pre-existing Febrile Illness. Febrile illness of any sort appears to predispose to heat illness¹¹,²⁰. Three men in this series had mild febrile symptoms within 24 hours of the exercise. None reported sick and excluding such highly motivated men from competitive events is difficult. Here education is vital. One man, who suffered severe heat stroke, was in the early stages of scrub typhus during the exercise and although feeling unwell felt unable to withdraw because of regimental pride.

Recent experience in Hong Kong has shown the continuing dangers to soldiers from high ambient temperatures. Much can be done to prevent heat illness by more extensive education of those exposed to the heat hazard and those supervising soldiers.

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REFERENCES
Heat Illness. A Report of 45 Cases from Hong Kong

A Henderson, J W Simon, W M Melia, J F Navein and B G Mackay

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