Effect of temperature on neuromuscular transmission in the Eaton-Lambert syndrome

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SUMMARY A patient with the Eaton-Lambert syndrome is described in whom no associated condition was discovered. There was clinical and electrical evidence that the defect in neuromuscular transmission became more severe as local temperature was raised.

The myasthenic syndrome of Eaton and Lambert is caused by a reduction in the number of acetylcholine quanta released at the neuromuscular junction (Elmqvist and Lambert, 1968), and is frequently but by no means always associated with a neoplasms (Brown and Johns, 1974). In the patient described, in whom no associated condition was discovered, the defect in neuromuscular transmission was influenced by local temperature. This finding has not previously been reported in the Eaton Lambert syndrome.

Case report

A 54 year old man was admitted to hospital in January 1978. Eighteen months earlier he had first noticed weakness and mild discomfort in the legs. Weakness progressed so that within four months he was no longer able to work as a ship's engineer. Deterioration in the legs was gradual from that time but there was in addition mild weakness of the arms. He had noticed that the weakness was considerably worsened by a hot bath or in sunny weather. Shortly before admission he had complained of tiring of the eyes and mild slurring of speech but there were no more definite bulbar symptoms and no sensory disturbances. His general health was good, and he had not smoked for 12 years.

On examination, abnormalities were confined to the neuromuscular system. There was mild weakness of neck flexion but no other cranial nerve abnormality. He had minimal weakness of upper limb muscles and more pronounced weakness of the trunk and of the proximal lower limb muscles, without wasting or fasciculation. He was unable to sit up from the supine position, and had a slow, somewhat waddling gait. Tendon reflexes were all either depressed or absent but could easily be elicited after 10 seconds of voluntary contraction of the muscle. Plantar reflexes were flexor. Sensation was normal.

Electrophysiological investigation showed the typical features of the Eaton-Lambert syndrome. Surface recording from the right abductor digit minimi muscle (ADM) showed the resting compound muscle action potential to be grossly reduced in amplitude, being 500 µV or less peak to peak. At all frequencies of repetitive stimulation there was an initial decrement in the amplitude of the resting muscle action potential. This was followed by no recovery at stimulus frequencies of 1–5/s; by recovery, often with subsequent increment, at 10/s; and by an increment of 200–1500% of resting values at higher (15–50/s) frequencies of stimulation. Stimulation after 10 seconds of maximum voluntary contraction of ADM resulted in action potentials of up to 5 mV. The right ulnar motor nerve conduction velocity in the forearm was 56 m/s; the right median and ulnar sensory nerve action potentials (finger to wrist) were 7.5 µV and 6.0 µV respectively. Electromyography of the right vastus medialis showed a “myopathic” pattern, with no spontaneous activity but many brief spiky motor unit potentials of small amplitude in a mildly reduced interference pattern.

Results of other investigations were normal, including full blood count, sedimentation rate, chest radiography, bronchoscopy, sputum cytology, serum urea, electrolytes including calcium and magnesium, thyroid, and liver function tests. After the patient’s report that warm conditions made
him weaker, further electrophysiological studies were carried out.

Methods

An active recording electrode was placed in a constant position over the belly of ADM, and an indifferent electrode was placed 40 mm distal to this point, over the tendon. The hand was immobilised in a splint and skin temperature was measured with a shielded thermistor probe on the thenar eminence. Local cooling was achieved by surrounding the wrist and hand with ice-filled bags, and an infrared lamp was used for warming.

Trains of stimuli of three seconds duration were delivered to the ulnar nerve at the wrist at 2/s and at 10/s, each after at least five minutes relaxation, and at 2/s immediately after 10 seconds maximal voluntary contraction of ADM. This procedure was carried out on several occasions at room temperature, after warming, and finally after cooling (skin temperatures respectively 33–35°C, 38–40°C, and 15–20°C). Two healthy subjects were used as controls.

Results

ACTION POTENTIAL IN RESTING MUSCLE

Over the temperature range 16–40°C there was a progressive decline in the compound muscle action potential. This effect was not seen in healthy subjects (Fig. 1).

POST-CONTRACTION POTENTIATION

With rising temperature there was a decline in compound muscle action potential when resting and after potentiation, but the amounts of potentiation increased. In healthy subjects these effects of temperature were not observed (Table).

<table>
<thead>
<tr>
<th>Table Effect of temperature on post-contraction potentiation</th>
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<tbody>
<tr>
<td>Cold</td>
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<td>Mean resting muscle action potential (N) (mV)</td>
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<tr>
<td>Mean post-contraction muscle action potential (C) (mV)</td>
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<td>Mean post-contraction potentiation (C/N)</td>
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STIMULATION AT 2/s AFTER REST

Decrement to about 50% of the initial muscle action potential, without recovery, occurred at all temperatures in the patient, but no decrement was observed in healthy subjects.

STIMULATION AT 2/s AFTER MAXIMAL VOLUNTARY CONTRACTION FOR 10 SECONDS

At room temperature and on warming, the initial (post-contraction) compound muscle action potential declined progressively during repetitive stimulation. On cooling, not only was the initial action potential larger than at higher temperatures but in addition repetitive stimulation resulted in a further small increase in amplitude, which reached a maximum at one second and was sustained. No effect was seen in healthy subjects.

STIMULATION AT 10/s AFTER REST

The effects of stimulation at 10/s are shown in Fig. 2. Ratios of successive muscle action potentials to the initial one are plotted. Serial mean values in the cold were larger than those at room temperature and the initial decrement was virtually abolished. On warming, the mean values were smaller than those at room temperature from 1.5 s onwards. In healthy subjects the maximum increment was 50% and occurred at room temperature.

Discussion

The effect of temperature in the Eaton-Lambert syndrome has not been described previously. In myasthenia gravis, now known to be predominantly
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Fig. 2  Repetitive stimulation at 10/s. Mean values of the ratio of successive compound muscle action potentials (MAP) to the initial (resting) potential, at 0.5 s intervals. Effect of temperature in the patient. ■ = cold (16–20°C); ▲ = room temperature (33–35°C); ● = warm (38–40°C).

A postsynaptic disorder, neuromuscular blockade is increased with increasing temperature (Borenstein and Desmedt, 1975).

The effects of temperature on several types of presynaptic blockade have been reported. In the rat nerve-muscle preparation the blockade induced by raising the ambient magnesium concentration is enhanced by raising the bath temperature over the range 20–40°C (Hubbard et al., 1971). Blockade induced by botulinum toxin is temperature dependent in the same direction (Wright, 1955) as is that produced by the tick I. holocyclus (Cooper and Spence, 1976). The present patient was clinically and electrophysiologically typical of the Eaton-Lambert syndrome which has close physiological similarities to the above presynaptic disorders. Rising temperature over the range 16–40°C caused a decline of up to 80% in the compound muscle action potential of both resting and post-exercise muscle. Over the same range there appeared to be some increase in post-contraction potentiation but the amount of contraction was not fully standardized. Moreover, only on cooling did repetitive stimulation cause further facilitation of the post-contraction muscle action potential. The most striking effect was the decline in response to stimulation at 10/s observed as the temperature was raised from 16–40°C. Rising temperature may be associated with a declining probability of acetylcholine release (Hubbard et al., 1971), and this is one possible explanation for the effect of temperature on neuromuscular transmission in the Eaton-Lambert syndrome.

Our findings have obvious practical implications for patients. In addition, they add to the acknowledged diagnostic difficulties in this condition (Brown and Johns, 1974). The mainly proximal distribution of muscle weakness, and the small spiky motor unit potentials of decreased mean duration typically found on electromyography (Rooke et al., 1960) may lead to the misdiagnosis of myopathy. Recording at low temperatures may result in misleadingly normal amplitudes of compound action potentials while the characteristic incremental response to repetitive stimulation is best seen in the cold and may indeed be missed at higher temperatures, particularly if rates above 10/s are not used. The most reliable diagnostic test remains that of stimulation after voluntary contraction (Elmqvist and Lambert, 1968), since the compound muscle action potential shows a large increment at all three temperature ranges.

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References


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