Muscle cramps may be caused by fluid and salt loss induced by diffuse or focal hyperhidrosis. Recent reports have described the efficacy of botulinum toxin in the treatment of primary focal hyperhidrosis. Botulinum toxin inhibits sweating by blocking exocytosis of acetylcholine from presynaptic cholinergic nerve terminals.

We report the case of a patient who complained of frequent muscle cramps associated with unusually severe axillary hyperhidrosis. We used botulinum toxin to treat the excessive focal sweating presuming that it would also reduce the muscle cramps. A total dose of 200 MU of botulinum A toxin (Dysport) per axilla markedly reduced sweating and cramps. The beneficial effect started four days after the injection and it was still present five months later. Treatment was repeated in the sixth month with analogous results. No side-effects were observed and no compensatory sweating occurred.

Key words: Cramps; axillary hyperhidrosis; botulinum toxin.

Introduction

Muscle cramps are common symptoms of neuromuscular disorders and are caused mostly by hyperactivity of the peripheral or central nervous system (Bertolasi et al., 1993; Layzer, 1994). The major causes of muscle cramps are lower motor neuron diseases, radiculopathies, metabolic disorders (i.e., hypothyroidism and hypoadrenalism) and extracellular volume depletion after diarrhea, vomiting, diuretic therapy and hemodialysis. An additional cause of cramps is the acute fluid and salt loss through excessive sweating (i.e., primary hyperhidrosis), probably owing to the increased mechanical effect of muscle fibers on the nerve terminals in a contracted extracellular space (Layzer, 1994). Primary hyperhidrosis is a disorder that generally involves the palms, soles or axillae. Its prevalence is 0.6 to 1.0% in young adults (Adar et al., 1977). It is an annoying complaint and causes affected patients severe social and psychological problems; conventional medical or surgical treatments produce only a partial benefit or are associated with potentially serious side effects (Malone et al., 1986; Byrne et al., 1990; Freeman et al. 1992; Drott et al., 1995; Bushara et al. 1996).

Recently, several reports have shown the efficacy of botulinum toxin in the treatment of palmar and axillary hyperhidrosis (Schinder et al. 1996; Naumann et al., 1997; Schinder et al., 1997; Shelley et al., 1998; Naumann et al., 1998; Odderson, 1998; Heckmann et al., 2001). Botulinum toxin owes its successful use in the treatment of this condition to its blocking action of sympathetic postganglionic fibers to sweat glands (Bushara and Park, 1994).

In this study, we report a patient who complained of diffuse muscle cramps and severe axillary hyperhidrosis treated with botulinum toxin.
using a compound containing alizarin red powder 25 g, rice starch 50 g, and dehydrated sodium carbonate 25 g. This formula produces an exceedingly volatile substance and does not adhere to dry skin but, when painted onto sweated skin, adheres readily and changes to an intense violet color.

A total dose of 200 MU (mouse unit) of botulinum A toxin (Dysport, Ipsen), diluted with 0.5 ml of sterile saline solution, was injected subdermally into the right axilla in ten different sites (20 MU per site), using a 0.33 mm gauge needle, after informed written consent from the patient and ethical committee approval. We injected low single doses of botulinum toxin (20 MU) diluted in 0.5 cc of saline solution instead of one cc as recommended by suppliers to distribute the toxin more uniformly in the area of excessive sweating without unnecessary spread to the surrounding regions. After one week, the left axilla was treated similarly.

Six months later, an analogous treatment was repeated in both axillae.

In the period following the treatment, the patient did not use drugs or change her way of life.

**Results**

Four days after receiving the first botulinum toxin injection, the patient noted that excessive sweating in the right axilla had already diminished. At the same time, the muscle cramps became less severe.

Two weeks after the treatment, the alizarin red test showed a marked reduction in sweating in most areas of the right axilla; only one small area (2 mm × 3 mm) of excessive sweating remained in a linear medial region. The patient no longer complained of muscle cramps.

Further assessment two weeks after injection of the left axilla again showed markedly reduced sweating (Fig. 1); the beneficial effect in the right axilla persisted and muscle cramps disappeared.

Botulinum toxin effect on sweating and muscle cramps was still present five months after the first injection. In the sixth month, the patient referred a slow and progressive reappearance of the hyperhidrosis which, later, became as severe as before treatment and was again associated with muscle cramps. At the end of the sixth month, we repeated botulinum toxin administration as previously described; results analogous to the first treatment were obtained.

No side-effects were reported. No compensatory sweating occurred.

**Discussion**

This report shows that muscle cramps associated with axillary hyperhidrosis can be effectively,
safely and persistently controlled by the local treatment of hyperhidrosis with botulinum toxin A. The treatment of severe benign muscle cramps is often difficult because systemic drugs may cause ill tolerated side-effects. A recent report has described successful treatment of muscle cramps by a neuromuscular block produced by local injection of botulinum toxin into the cramping muscle (Bertolasi et al., 1997). In our study, we now show that botulinum toxin is also effective in treating muscle cramps secondary to hyperhidrosis by blocking the excessive sweating. In explaining why the muscle cramps disappeared, we think a direct reduction by spread of botulinum toxin through the bloodstream unlikely. Indeed, in a preceding study, after injection of botulinum toxin into the calf muscles of patients who had a benign cramp-fasciculation syndrome, the cramps on the treated side disappeared but the cramps on the untreated side remained (Bertolasi et al., 1997). Furthermore, the patient we now describe never had muscle weakness or other side effects linked to the action of botulinum toxin on motor endplates.

In conclusion, our case confirms that acute fluid and salt loss may cause muscle cramps and suggests that botulinum toxin, although it needs repeated administration over time (every five months), may be effective in treating primary axillary hyperhidrosis and may also resolve the co-existing muscle cramps.

REFERENCES


