A Case of Painful Hemimasticatory Spasm with Masseter Muscle Hypertrophy Responsive to Botulinum Toxin

Jin-Hyuck Kim a
Seok-Won Han b
Yun Joong Kim b
Jooyong Kim b
Mi-Suh Oh b
Hyeo-Il Ma b
Byung-Chul Lee b

aDepartment of Neurology, Keyo Medical Foundation Keyo Hospital, Uiwang, Korea
bDepartment of Neurology, Hallym University, College of Medicine, Anyang, Korea

Hemimasticatory spasm (HMS) is a rare disorder of the trigeminal nerve characterized by paroxysmal involuntary contractions of the unilateral jaw-closing muscles. HMS has been frequently described in association with facial hemiatrophy or localized scleroderma, but hypertrophy of an involved muscle may occur without these findings; few cases have been associated only with facial hemihypertrophy. In this paper, we present a case involving HMS associated with marked hypertrophy of the left masseter and relate the electrophysiological, radiological, and clinical findings. This patient was successfully treated with repeated local injections of botulinum toxin.

Case Report

A 42-year-old female presented with involuntary paroxysmal spasms of the left masticatory muscle, of 6 months duration. The spasms were mild in the early stage; however, they increased gradually in duration and frequency. The spasms were frequently painful and sometimes awoke her from sleep. The lower half of the face on the left side hypertrophied over time. She had no specific medical diseases.

On admission, her masseter muscle on the left was markedly hypertrophied. However, neither FHA nor a skin lesion was noticed. The spasms usually continued for about 10 seconds, but sometimes lasted for up to a few minutes. Neurologic examinations, including masticatory muscle function and jaw jerk, were normal. Hematological, biochemical, and serologic tests, including antinuclear antibody, rheumatoid factor, anti-DNA antibody, and C-reactive protein, were normal. The blink reflex was normal on an electrophysiological study. Masseter reflexes elicited by tapping her chin and recorded by surface electrodes from the masseter muscle showed delayed latencies and decreased amplitudes on the left side during the spasm episodes (Figure 1A). Moreover, the masseteric silent period was attenuated on the affected side (Figure 1B). Surface electromyography demonstrated irregular bursts of motor unit potentials at 50 to 200 Hz during the period of involuntary spasm or at random intervals (Figure 1C). Magnetic resonance imaging of the head showed pro-
nounced hypertrophy of the left masseter muscle (Figure 2). A muscle biopsy done at the central region of the hypertrophied muscle revealed nonspecific results.

The frequency of the spasm and pain improved slightly after treatment with phenytoin. Several local injections of 20-50 units of botulinum toxin in the left masseter muscle led to marked improvement of her symptoms. After 3 months, the hypertrophy of the masseter muscle was substantially decreased (Figure 3). The benefits of the botulinum toxin injections persisted for 4 months, and subsequent injections were performed with equal success. The follow-up electrophysiological profiles were also improved.

**Discussion**

Since the first reported description of the electrophysiological and clinical features of HMS with FHA,1 about 20 cases have been described. HMS predominantly affects women in the third to fourth decades and is associated with hemiatrophy in two-thirds and scleroderma in one-third.2–4 The spasm affects the masseter muscle most frequently and may involve one or more jaw-closing muscles.2,3,8,9 However, the involvement of jaw openers has never been described.2,3

Although the pathophysiology of HMS is not well known, peripheral lesion of the trigeminal motor nerve is considered to play a role.2–7 It has been proposed that HMS is produced by ectopic discharge secondary to focal demyelination of the trigeminal motor fiber caused by compression, entrapment, or stretching injury to the extracranial portion of the nerve.2,5,8,9 Some authors have postulated that HMS might originate from vascular compression of the trigeminal motor nucleus or the motor root near the brainstem.2,3,5,9 However, evidence of vascular compression has not been demonstrated in surgical explorations.2,5

Medications such as carbamazepine, phenytoin, and clonazepam may be helpful, although the effect of treatment with these drugs has been insufficient in most patients with HMS.3,5,7,8 In our case, phenytoin was beneficial to only a
limited degree for improving pain and the frequency of the spasms. On the other hand, the local injections of botulinum toxin resolved the spasms and reduced the muscle hypertrophy. In our search of the literature, another eight patients with HMS received treatment with botulinum toxin, and all of them showed an excellent response to this therapy (Table 1).2-5,7-9 All of the cases except two7,9 were associated with muscle hypertrophy. The reports all described obtaining marked improvements in the pain and spasms; however, an effect on the muscle hypertrophy was not mentioned in the previous cases. Our case clearly demonstrated a regression in the size of the hypertrophied muscle.

In this study, we presented a case of HMS associated with masseter muscle hypertrophy and related the electrophysiological and neuroimaging findings. Botulinum toxin may be considered the treatment of choice for pain and spasm in HMS. In addition, this therapy can be helpful for decreasing the size of the hypertrophied muscle.

REFERENCES