INTRODUCTION

Microvascular decompression (MVD) has been known to be a very effective and relatively safe surgical modality in the treatment of hemifacial spasm. But rare debilitating complications have been reported such as cranial nerve dysfunctions. We have experienced a very rare case of unilateral soft palate palsy without the involvement of vocal cord following microvascular decompression. A 33-year-old female presented to our out-patient clinic with a history of left hemifacial spasm for 5 years. On postoperative 5th day, patient started to exhibit hoarseness with swallowing difficulty. Symptoms persisted despite rehabilitation. Various laboratory work up with magnetic resonance image showed no abnormal lesions. Two years after surgery patient showed complete recovery of unilateral soft palate palsy. Various etiologies of unilateral soft palate palsy are reviewed as the treatment and prognosis differs greatly on the cause. Although rare, it is important to keep in mind that such complication could occur after microvascular decompression.

Key Words : Microvascular decompression · Hemifacial spasm · Lower cranial nerve · Soft palate palsy · Vocal cord.

CASE REPORT

A 33-year-old female was presented to our out-patient clinic with a history of left hemifacial spasm for 5 years. Her past medical history was unremarkable. The patient underwent microvascular decompression in 2010 at our medical center (Fig. 1).

Intra-operatively, bilateral facial electromyography and brainstem auditory evoked potential was monitored. After the dura incision, gentle retraction of the cerebellum was applied using Greenburg retractor and we have exposed the lower cranial
nerves. Dissection of the arachnoid was done from the rostral border of the lower cranial nerves. The arachnoid band was very tough and difficult to dissect, but no obvious injury occurred to the lower cranial nerve during the dissection. The left anterior inferior cerebellar artery was the offending vessel compressing the facial nerve from the caudal side and decompression was successfully completed without any adverse events and with no change in the intraoperative monitoring of brainstem auditory evoked potential. Patient recovered well from anesthesia with reduced facial spasm, compared to the pre-operative state. Postoperatively, neurologic exam revealed no abnormalities and also patient’s swallowing and vocalization function was intact. Immediate postoperative brain CT scan performed on postoperative day showed no sign of hemorrhage.

On postoperative 5th day, the patient started to exhibit hoarseness with swallowing difficulty. Barium swallowing test was done to evaluate dysphagia, and minimal nasal regurgitation was demonstrated due to unilateral soft palate palsy. On endoscopic exam, performed by an otolaryngologist, the vocal cord was intact in both sides with unilateral left soft palate palsy, which suggested tensor and/or levator veli palatini muscle weakness (Fig. 2). Under the impression of possible lower cranial nerve palsy caused by postoperative edema of the nerve, steroid (prednisolone) was prescribed and we decided to observe the symptoms. Symptoms persisted with slight improvement with steroid therapy but repeatedly aggravated with steroid tapering. She received speech rehabilitation therapy for persistent difficulty of phonation and swallowing.

On postoperative 19th day, patient complained of difficulty breathing during sleep. Polysomnography was done to evaluate sleep apnea. Mild obstructive sleep apnea syndrome was observed secondary due to soft palate palsy. We encouraged her to sleep on her side; then her symptom improved.

Despite persistent speech rehabilitation, symptoms still persisted. We performed cranial nerve MRI to evaluate for the possibility of any structural abnormalities, such as delayed hematoma or possible infection. However, MRI showed no abnormal structural lesions or abnormal enhancement, which may suggest infection (Fig. 3). The patient continued to receive speech rehabilitation with subjective improvement of swallowing and speech.

On postoperative 25th day, the patient developed grade II facial palsy by House-Brackmann grade on the left side. Patient's facial palsy improved with administration of acyclovir for 5 days, but hoarseness with swallowing difficulty showed little improvement.

The patient received prolonged speech and swallowing rehabilitation therapy, and follow up examination was done before discharge showed improvement of nasalization, but persistent hypernasality. Patient was discharged with recommendation for prolonged rehabilitation.

Patient visited our out-patient for a follow up in postoperative 2 years. Endoscopic examination revealed normalized soft palate palsy and no abnormality in the vocal cord mobility (Fig. 4). However, patient's subjective difficulty of speech still persisted.
and we decided to observe her symptoms for any changes in the future.

DISCUSSION

Cranial nerve palsy has been reported as complication of MVD and can cause symptoms that markedly decrease the quality of life. Most commonly, the 7th and 8th cranial nerve are involved with rare lower cranial nerve involvement. Among the lower cranial nerves, glossopharyngeal nerve and vagus nerve are most commonly involved.

Glossopharyngeal nerve contains both motor and sensory branches that supply the tongue and stylopharyngeus muscle of the pharynx. Impairment of glossopharyngeal nerve would cause decreased sensation in the pharynx including taste in the posterior one third of the tongue with difficulty in swallowing and dysphonia\(^{2-4}\). Our patient developed swallowing difficulty with dysphonia, which suggests possible dysfunction of the 9th nerve.

Vagus nerve also contains both motor and sensory branches that innervate all striated muscles of the laryngopharynx, except the stylopharyngeus and tensor veli palatini. Patients with unilateral high vagal lesions would suffer from breathy dysphonia and aspiration because of impaired vocal cord movement. Due to the loss of sensation of the laryngopharynx, causing loss of gag reflex, patients might also suffer severe dysphagia related to the poor coordination of the oro- and hypo-pharyngeal phases of swallowing\(^{2-4}\). Unilateral soft palate palsy with dysphonia and dysphagia was observed in our patient, which suggests unilateral vagus nerve dysfunction, but the sparing of the vocal cord suggest the function of the vagus nerve is only partially impaired.

Although the functions and innervations of the 9th and 10th nerve are known, it is difficult to exactly locate which lower cranial nerve is causing the symptom by physical examination. Many symptoms are subtle and overlap with other diseases, making the diagnosis elusive\(^3\).

Common injury to the vagus nerve is accompanied by vocal fold palsy, but our patient only presented with soft palate palsy. In addition, partial injury of the vagus nerve with possible involvement of the glossopharyngeal nerve have never been reported as a complication of MVD in prior literature, thus making it difficult to identify the exact cause of such peculiar symptoms.

Several etiologies are reported in literature that can cause lower cranial nerve paralysis. Skull base tumors are the most common cause and trauma or surgery above the skull base and from the cerebellopontine angle can also be a cause. Rarely, infection and ischemic vascular insult are reported to cause unilateral vagus nerve palsy\(^{3,7,8,12}\).

It is important to note that in our patient, the hoarsness and dysphagia developed several days after the surgery. Generally, lower cranial palsy caused by trauma would occur immediately after the event, which implies the possibility of other factors, such as infection contributing in the delayed onset of lower cranial nerve palsy.

In the literature, some have reported herpes simplex virus (HSV) to be a possible cause of unilateral vagus nerve palsy\(^{12}\). In addition, varicella zoster virus is also reported to cause lower cranial nerve palsy\(^3\). Although we have failed to show such connection in our patient, history of delayed facial palsy also suggests the possibility of concurrent viral infection. Others have also suggested that delayed facial palsy after MVD in patients with HFS may be caused by viral infections. As HSV is widely known to cause Bell’s palsy, it could likewise cause reversible inflammation of vagus nerve causing such clinical presentations\(^{11}\).

The history of relapsing symptom, followed by tapering of steroids, suggests additional evidence that inflammatory reaction contributed in the development of such symptom. Antiviral agent, such as acyclovir, is reported to be an effective treatment that we have used for facial palsy of the patient.

Another possibility can be suggested. Some reported rare anatomical variants of lower cranial nerves with interconnection between the glossopharyngeal nerve and vagus nerve. Such interconnection is so thin and very fragile that it makes them prone to shearing stress\(^{14}\). Though they have not suggested its clinical implication, it is possible that such interconnection holds some significance in the nerve conduction. As such, it may have been overlooked and damaged during the retraction of the surgery, causing lower cranial nerve palsy.

The clinical course of unilateral lower cranial nerve palsy is variable depending on the cause. Initial extensive laboratory work up with CT and MR imaging is necessary to evaluate the cause. If no particular structural abnormalities are noted, conservative treatment, including acyclovir and steroid administration with speech therapy, seems to be reasonable. If symptoms persist, surgical treatments, such as medialization laryngoplasty and laryngeal reinnervation, can be considered\(^{8,13}\).

We have experienced a very rare case of unilateral soft palate palsy, following MVD. Although the exact cause of this complication had not been identified, we have suggested possible causes for such rare complication. It is important to keep in mind that MVD is performed to improve the quality of life rather than to save life. Although it has relatively little complications, rare and debilitating complications must not be overlooked, and caution must be taken to avoid them.

CONCLUSION

MVD is a very effective and relatively safe surgical modality in the treatment of HFS. However, rare debilitating complications are reported, such as cranial nerve dysfunctions. The lower cranial nerve palsy we are reporting is one of such complication that can markedly decrease the quality of life of a patient and it is important to note that such delayed complication can occur without evident injury to the nerve during the operation. It is important to consider various etiologies as the treatment, and prognosis differs greatly on the cause. Additionally, careful surgical technique with minimal retraction with extra care to avoid
damage to the lower cranial nerve during exploration may be important to minimize such complication.

References