Trochlear Nerve Schwannoma with Intratumoral Hemorrhage Presenting with Persistent Hiccups: A Case Report

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Abstract

Trochlear nerve schwannoma without neurofibromatosis is extremely rare. To our knowledge, only 31 surgical cases have been reported to date, and only 2 cases of trochlear nerve schwannoma with intratumoral hemorrhage have been reported. None of those cases presented with persistent hiccups. We report the case of a 44-year-old man with trochlear nerve schwannoma associated with intratumoral hemorrhage who presented with a 10-day history of persistent hiccups. Computed tomography and magnetic resonance imaging revealed a solid tumor with a 3-cm diameter and intratumoral hemorrhage in the left petroclival region that compressed the midbrain andpons. Subtotal removal of the tumor was performed via the zygomatic transtemporal approach. Intraoperative findings revealed a tumor arising from the trochlear nerve. The histologic diagnosis was schwannoma of Antoni type A cells with intratumoral hemorrhage. Although the patient’s left trochlear nerve palsy worsened temporarily, his postoperative course was uneventful. We present this rare case and discuss the mechanism underlying the patient’s persistent hiccups.

Keywords

► persistent hiccups
► trochlear nerve schwannoma
► intratumoral hemorrhage

Introduction

Schwannomas are tumors originating from Schwann cells and account for 8% of all primary brain tumors.1 Although schwannomas develop mainly in the sensory nerves, most often the vestibular nerve, in very rare cases they can develop in the trochlear nerve, which is a motor nerve.1,2 To our knowledge, only 31 cases of trochlear nerve schwannoma have been reported.3–32 In many of these cases, the patient initially presented with headache, double vision (associated with trochlear nerve paralysis or oculomotor paralysis), hemiplegia, or cerebellar symptoms; however, none of the patients had an initial presentation of persistent hiccups. Furthermore, a schwannoma complicated by intratumoral hemorrhage is rare, with only 2 of the 31 reported cases of trochlear nerve schwannoma exhibiting this complication.23,31

We report a very rare case of trochlear nerve schwannoma with an intratumoral hemorrhage that was the likely trigger of persistent hiccups and was treated with tumor resection.

Case Report

A 44-year-old man was referred to our hospital because of persistent hiccups lasting for 10 days. The patient did not complain of double vision, although his results for the Bielschowsky head-tilt test were positive, which suggested left trochlear nerve paralysis. There were no signs of disorders
affecting the cerebral nerves including the oculomotor, trigeminal, and abducens nerves. The patient also showed no signs of hemiplegia, cerebellar symptoms, or headache. The hiccups subsided spontaneously on the day following the patient’s first visit to our facility.

Computed tomography (CT) revealed the presence of an isodense tumor, which was partially accompanied by a hyperdense inner region, in the left petroclival area (Fig. 1A). Magnetic resonance imaging (MRI) also revealed a well-demarcated tumor (27 × 27 × 30 mm) in the left petroclival area. The tumor had mixed intensity (isointensity to hypointensity) on a T1-weighted image, mixed intensity (isointensity to hyperintensity) on a T2-weighted image, isointensity on a fluid-attenuated inversion-recovery image, and hypointensity (consistent with the high-intensity area on the CT scan) on a T2*-weighted image (Fig. 1B). The tumor showed heterogeneous contrast enhancement in the presence of the contrast agent gadolinium (Fig. 1C, D). Imaging revealed that the tumor compressed the midbrain and pons from the left anterior direction, although there was no significant change of intensity in these regions. A heavy T2-weighted image showed that the tumor was in contact with the left oculomotor nerve in the superior region and with the left trigeminal nerve in the inferolateral region.

Tumor resection was performed using the zygomatic transpetrosal approach (consisting of an anterior transpetrosal approach and a zygomatic arch resection). Although this allowed the tumor to be freed from the trigeminal nerve, the trochlear nerve had become partially assimilated by the tumor membrane, leading to the diagnosis of a trochlear nerve-derived tumor (Fig. 2A). Because the tumor was partially but strongly attached to the brainstem and clivus, its membrane was incised for internal decompression. A blood clot found inside the tumor, implying intratumoral bleeding, was then removed (Fig. 2B). The tumor was resected while keeping the tumor membrane close to the brainstem and clivus and while leaving the medial membrane intact. Because a sizable air cell had developed at the petrous bone, we collected fat from the abdominal region, and after
bone resection, the defect was filled with fat to complete the operation. Postoperative histopathologic examination revealed entangled and proliferating spindle cells with partially swollen quasi-circular nuclei, which were indicative of a schwannoma composed primarily of Antoni type A cells (Fig. 3A). Traces of bleeding accompanied by hyalinized blood vessels and hemosiderin deposition were noted inside the tumor (Fig. 3A inset), and the removed blood clot contained blood from both recent and previous bleeding (Fig. 3B).

Postoperative MRI confirmed that the tumor had been removed while keeping only the left membrane intact (Fig. 4). The patient showed temporary postoperative aggravation of left trochlear nerve paralysis, but no new symptoms appeared. Since then (1 year has passed), the symptoms have not recurred. In addition, MRI performed 6 months after the operation showed no change in the size of the residual tumor.

**Discussion**

- Table 1 summarizes the 31 surgical cases that have been reported to date, together with the case reported here (3–32). Patients with trochlear nerve schwannoma often present initial symptoms of double vision (60%), hemiplegia (43%), headache (40%), and cerebellar signs (37%) (Table 1). Although cases of this tumor presenting with diverse symptoms have been reported, no case with persistent hiccups as the initial symptom was previously described.
Hiccups usually subside within a short time; those lasting ≥ 48 hours are defined as persistent hiccups, whereas those lasting ≥ 1 month are termed intractable hiccups. Gastric flatulence is generally responsible for hiccups, but persistent or intractable hiccups suggest a possible complication by an organic disease. Although the exact mechanism of hiccupping remains unclear, the hiccup reflex arc is known to be composed of several neural pathways. The afferent pathway involves the sensory fibers of the vagus nerve, the glossopharyngeal branch of the glossopharyngeal nerve, the pharyngeal plexus (C2–C4), and the sympathetic nerves (T6–T12). Its efferent pathway involves the glottis and phrenic nerve, which is linked to the auxiliary respiratory muscles. Linkage to the central nervous system has not been elucidated in detail, but possible interactions with the medulla oblongata (including the solitary nucleus and ambiguous nucleus), part of the pons, reticular formation of the brainstem, phrenic nerve nuclei, and hypothalamus have been suggested.

Causes of persistent or intractable hiccups originating in the central nervous system include vascular, infectious, and structural processes, which are thought to suppress the hiccup reflex. Furthermore, some case reports have attributed hiccups to brainstem lesions, particularly those in the medulla oblongata. In the present case, although persistent hiccups were observed as the initial symptom, after which they spontaneously subsided, the medulla oblongata had not been directly compressed by the tumor. This finding, together with the observation of brainstem compression by the tumor, suggests that the hiccups in this case may have been caused by intratumoral hemorrhage.

Schwannomas are only rarely complicated by intratumoral hemorrhage. Historically, clinically significant intratumoral hemorrhage was reported in only a small fraction of vestibular schwannomas. However, advances in imaging and larger analyses suggest that intratumoral hemorrhage is far more common than previously believed and might represent an aspect of the natural history of vestibular schwannomas. Hemorrhagic vestibular schwannomas often manifest with acute symptom onset and have been studied inadequately since McCoyd’s initial description. Of the 31 reported cases of trochlear nerve schwannoma, only 2 had intratumoral hemorrhage, and both these cases showed acute aggravation of symptoms (Table 2). According to a report published by Yamamoto et al, the initial symptoms of this condition include sudden headache, nausea, vomiting, and double vision. Ohba et al reported a case in which intratumoral hemorrhage occurred during the follow-up of a trochlear nerve schwannoma. Their case showed aggravation of the original symptoms (left hemiparesis, right trochlear nerve paralysis, and sensory disorder of the third branch region of the right trigeminal nerve), as well as developed right oculomotor paralysis and right facial palsy.

Table 1 Summary of 32 surgical cases of trochlear neurinoma including our case

<table>
<thead>
<tr>
<th>Study</th>
<th>Age (yrs)</th>
<th>Symptoms</th>
<th>Duration of symptoms</th>
<th>Size (mm)</th>
<th>Residual symptom</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yamamoto et al</td>
<td>37/F</td>
<td>Sudden onset of headache, nausea, vomiting, diplopia</td>
<td>2 wk</td>
<td>10 × 9 × 7</td>
<td>Right fourth palsy</td>
<td>5 y</td>
</tr>
<tr>
<td>Ohba et al</td>
<td>48/M</td>
<td>Diplopia, left hemiparesis, hypesthesia of the right face, right facial palsy</td>
<td>3 wk</td>
<td>25</td>
<td>Right fourth palsy</td>
<td>4 mo</td>
</tr>
<tr>
<td>Hatae et al (current study)</td>
<td>44/M</td>
<td>Persistent hiccup, left fourth palsy</td>
<td>10 d</td>
<td>27 × 27 × 30</td>
<td>Left fourth palsy</td>
<td>1 y</td>
</tr>
</tbody>
</table>

Table 2 Summary of three surgical cases of trochlear neurinoma with intratumoral hemorrhage
In the present case, intratumoral hemorrhage was confirmed by intraoperative and pathologic findings. CT performed during the first examination revealed interposition of hyperintense areas within the tumor, and an MRI of the same region revealed low signal intensity on T2*-weighted image. CT performed 2 weeks later revealed the presence of iso-intensity. These findings suggest that the patient had developed intratumoral bleeding several days before the first visit. In view of the patient's disease history, the hiccups probably began immediately after intratumoral bleeding, suggesting that this may have been their cause. The area affected by the hematoma was located caudal to the tumor. Considering that this may have been their cause, the spontaneous disappearance of the hiccups probably began immediately after intratumoral bleeding, leading to the onset of persistent hiccups. The lack of hiccups prior to intratumoral hemorrhage, as well as following surgery, supports this hypothesis. The spontaneous disappearance of the hiccups also indicates the involvement of indirect, rather than direct, stimulation.

Conclusions

We reported a case of trochlear nerve schwannoma in which intratumoral hemorrhage probably caused persistent hiccups. To our knowledge, this is only the third reported case of trochlear nerve schwannoma complicated by intratumoral hemorrhage, and the first case exhibiting persistent hiccups as the initial symptom.

Conflict of Interest

The authors have no personal financial or institutional interest in any of the drugs, materials, or devices in the article.

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