Isolated Hypoglossal Nerve Palsy Due to an Osteophyte with Atlantoaxial Dislocation

Ko Ozaki,1 Iwao Yamakami,2 Yoshinori Higuchi,3 and Toshio Fukutake4

Introduction

Hypoglossal nerve palsy is mostly caused by tumors and trauma in the craniovertebral junction and is commonly associated with either other cranial nerve palsies or corticospinal signs. Isolated hypoglossal nerve palsy (IHP), or hypoglossal nerve palsy without any other neurological signs, is rare. Atlantoaxial dislocation (AAD) is a loss of stability between the atlas and axis (C1-C2) and can occur in any age group from various causes including trauma, inflammation, congenital abnormalities, or unknown origins. AAD presents various clinical features ranging from neck pain to fatal outcomes, and the instability of C1-C2 often forms an osteophyte. We report an AAD patient who presented with left IHP due to hypoglossal nerve compression in the hypoglossal canal by an atlanto-occipital joint osteophyte.

Case Report

A 77-year-old woman presented with left neck pain for 1 month, followed by speech difficulties and the feeling of a swollen tongue on the left side for 3 days. She had been diagnosed with AAD without rheumatoid arthritis (RA) 2 years prior, and had no history of trauma (Fig. 1). Her medical history was not significant, except for hypertension. Physical examination showed a mild tongue deviation to the left when it extended. The tongue movement to the right was impaired, though the movement to the left was not. Her soft palate movement was normal with no hoarseness, swallowing disturbances, or other neurological signs. The neurological diagnosis was left IHP. A computed tomography (CT) scan showed the AAD and moreover, that an osteophyte involving the atlanto-occipital joint obstructed the external orifice of the left hypoglossal canal (Fig. 2). Magnetic resonance imaging (MRI) showed no mass lesions either intracranially or extracranially along the hypoglossal

Isolated hypoglossal nerve palsy (IHP), or hypoglossal nerve palsy without any other neurological signs, is rare. We report a woman with atlantoaxial dislocation (AAD) who presented with IHP due to hypoglossal nerve compression by an osteophyte at the hypoglossal canal. A 77-year-old woman presented with speech difficulties and the feeling of a swollen tongue on the left side for 3 days. Her only neurological feature was left hypoglossal nerve palsy. She had been diagnosed with AAD 2 years before. Computed tomography (CT) and high-resolution magnetic resonance imaging (MRI) revealed the compression of the basicranial hypoglossal nerve at the external orifice of the hypoglossal canal by an AAD osteophyte which was causing IHP. IHP can develop due to hypoglossal nerve compression by an osteophyte from AAD. CT and high-resolution MRI revealed this rare mechanism of IHP.

Keywords: hypoglossal nerve, atlantoaxial dislocation, craniovertebral junction, cranial nerve palsy

1Narita Red Cross Hospital, Department of Neurosurgery, Narita, Chiba, Japan
2Seikeikai Chiba Medical Center, Department of Neurosurgery, Chiba, Chiba, Japan
3Chiba University Graduate School of Medicine, Department of Neurological Surgery, Chiba, Chiba, Japan
4Kameda Medical Center, Department of Neurology, Kamogawa, Chiba, Japan

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Fig. 1 Abnormal atlantodental interval revealed by sagittal bone CT. CT: computed tomography.
nerve. High-resolution MRI (balanced fast field echo; Philips Healthcare, Best, Netherlands) revealed a discontinuity in the basicranial segment of the left hypoglossal nerve at the external orifice of the hypoglossal canal (Fig. 3). The radiologic diagnosis was IHP due to AAD osteophyte compression of the hypoglossal nerve. Surgical intervention to decompress the hypoglossal nerve was not performed after weighing its invasiveness and effectiveness. Her symptoms and signs were stable during the 1-year follow-up period.

Discussion

The trajectory of the hypoglossal nerve, from the brainstem to the lingual muscles, comprises five segments: medullary (nuclear), cisternal, basicranial, retro- and prestyloid, and submandibular. Intracranially, the hypoglossal nerve arises from the hypoglossal nucleus in the medulla, travels along the dorsal side of the vertebral artery, penetrates the dura mater, and runs into the hypoglossal canal. Extracranially, this nerve runs near the internal carotid artery, the vagus and accessory nerves, and communicates to the vagus, first, and second cervical nerves. Damage to any point along this trajectory from the medulla to the lingual muscles causes a hypoglossal nerve palsy, and in the present case, the compression of the hypoglossal nerve at the hypoglossal canal by an osteophyte caused hypoglossal nerve palsy. Because of the anatomical characteristics of the hypoglossal nerve, hypoglossal nerve palsy is usually associated with other cranial nerve palsies (IX, X, and XI) and/or pyramidal signs, and IHP is rare.

The most frequent cause of IHP is skull base tumors including metastases, meningiomas, and malignant tumors. Less frequently, IHP develops due to trauma, vascular pathologies, and inflammatory etiologies, among others. Except for the present case, two cases of IHP due to osteophytes have been reported. One case was of a 92-year-old woman without any AAD history; the other case was of an 86-year-old woman with a history of cervical spondylotic myelopathy requiring C2-4 laminectomy 2 years previously. Both cases were of elderly patients with no AAD history. This is the first case report of IHP due to an osteophyte from AAD. AAD resulted in the development of an osteophyte involving the atlanto-occipital joint, which resulted in mechanical compression of the basicranial segment of the hypoglossal nerve in the hypoglossal canal. RA often leads to AAD. Hypoglossal nerve palsy caused by AAD with RA was bilateral in the three cases which Blankenship et al. reported. The unilateral IHP in the present case was caused by an osteophyte without RA.

Diagnosing the exact mechanism of IHP is often challenging. In the present case, CT and high-resolution MRI diagnosed the osteophyte compression of the basicranial segment of the hypoglossal nerve in the external orifice of the hypoglossal canal.

The hypoglossal canal area can be approached surgically via either a transnasal, transoral, or a transcranial approach. At present, any approach to the hypoglossal canal is associated with considerable invasiveness and major risk and may not lead to recovery from hypoglossal palsy by the surgical removal of an osteophyte. Further advances in surgical techniques may be more effective and less invasive in decompressing the hypoglossal nerve at the hypoglossal canal. Local steroid injection is recommended as a non-surgical treatment for the entrapment neuropathy. However, the effect is transient and local steroid injection into the hypoglossal canal is difficult and dangerous because of anatomical characteristics of the canal.
**Conclusion**

This is the first case report of IHP due to AAD osteophyte compression of the hypoglossal nerve at the hypoglossal canal, which was diagnosed by CT and high-resolution MRI.

**Conflicts of Interest Disclosure**

The authors declare no conflicts of interest associated with this manuscript.
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Corresponding author:
Ko Ozaki, MD, Department of Neurological Surgery, Chiba University Graduate School of Medicine, Inohana 1-8-1, Chuo-ku, Chiba, Chiba 260-8670, Japan.
✉️ ozako.1115@gmail.com