Surgical Treatment of Rotational Vertebral Artery Syndrome Induced by Spinal Tumor: A Case Report and Literature Review

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Vertebrobasilar insufficiency (VBI) provoked by physiological head rotation is known as rotational vertebral artery syndrome (RVAS) or Bow Hunter syndrome. RVAS most often occurs at C1–2 level with head rotation and presents with symptoms of VBI. Several previously published studies have reported RVAS at subaxial sites (V2 segment), however, tumor-induced RVAS has never been reported. The authors report the first case of RVAS at V2 segment due to compression from a spinal tumor. A 71-year-old man presented with symptoms of dizziness provoked by head rotation or neck extension. Computed tomography (CT) angiography and dynamic cerebral angiography revealed circumferential stenosis with neutral neck position and complete occlusion of the left dominant vertebral artery (VA) at C5 level with his neck extended or rotated to the left. Complete neurological recovery was achieved after removal of a spinal osteochondroma and surgical decompression of the left VA via an anterior approach. Spinal tumors should be included in the differential diagnosis in cases of RVAS. Spinal degenerations or sarcomatous transformation of the tumor could lead to clinical manifestations of RVAS in cases with spinal osteochondroma. Complete removal of the tumor with or without spinal fusion would be the treatment of choice, in addition to medical treatment in the cases of acute stroke.

Keywords: rotational vertebral artery syndrome, vertebrobasilar insufficiency, spinal osteochondroma, subaxial spine, anterior approach

Introduction

Rotational vertebral artery syndrome (RVAS), also known as Bow Hunter syndrome, is the clinical presentation of vertebrobasilar insufficiency (VBI) produced by reversible occlusion of the vertebral artery (VA) during head rotation.

Typically, the dominant VA is transiently compressed at the C1-2 level, but there are multiple reports indicating that RVAS can occur in the subaxial cervical spine. The majority of these cases were secondary to spondylotic changes, such as osteophytes of the uncovertebral joint, disc herniation, and spondylolisthesis.1-5 Here, we report a rare case of RVAS caused by a compressive spinal osteochondroma that occurred at the C5 transverse foramen. To our knowledge, a RVAS case triggered by spinal tumor has not been previously reported. The case was successfully treated with total removal of the tumor via an anterior approach. We discuss the surgical and pathological findings of this rare entity in detail.

Case Report

A 71-year-old male presented to Nagoya University Hospital complaining of a 2-year history of dizziness, without loss of consciousness, provoked by neck extension or rotation of his head nearly 90° toward the left. The symptom always resolved immediately after he returned his head to the neutral position. He had never suffered from any pathology of the cervical spine and had a negative family history. Neurological examination was otherwise normal without myelopathy, motor weakness or sensory disturbance.

Cervical spine x-rays showed degenerative change with collapsed disc space (Fig. 1A) and segmental hypermobility of the index level on flexion and extension X-ray (Figs. 1B, and 1C) compared with other levels, while there was not significant spinal instability. Computed tomographic angiography (CTA) of the VA obtained with his head maintained in neutral position showed severe circumferential stenosis of the left VA with medial displacement by an exophytic bony structure as well as with lateral intervertebral disc bulging at the C5 transverse foramen (Figs. 2A, 2B, and 2D–2G). The bony mass was presumed to be arising from the C5/6 facet joint on computed tomography (CT) (Figs. 2A, 2B, and 2D). Spinal degenerations, such as lateral disc bulging, multilevel facet joint osteoarthritis and osteophyte formation, were also confirmed on CT and magnetic resonance image (MRI) (Figs. 2C, 2F, and 2G). Cerebral angiography demonstrated stenosis of the left VA at the C5/6 level with his head maintained neutral (Fig. 3A) and a right VA terminating at the posterior inferior cerebellar artery (PICA) (Fig. 3C). However, on dynamic cerebral angiography with his head rotated to the left or extended, the left VA was completely occluded and the distal VA flow was weakly filled by collateral circulation from muscular branches (Fig. 3B).
Surgery was recommended to remove the bony mass and decompress the left VA. A left-sided anterior approach was adopted. The longus colli muscle was dissected off the vertebral body at the C4 to C6 levels on the left side and mobilized laterally. The muscle was divided transversely at the index level with the fascia of the muscle left intact to preserve sympathetic nerves. The anterior surface of the C5 transverse process was exposed, then unroofed. Cartilage-like solid tissue was noted inside the transverse foramen, compressing the VA towards the medial direction (Fig. 4A). The lesion was adherent slightly to the VA at the medial aspect and heavily to the C5/6 facet joint, therefore, it was gently separated circumferentially and successfully removed en bloc. VA nicely expanded and pulsated immediately after removal of the mass lesion. Medial aspect of VA was not explored since we were concerned additional maneuver of severely squeezed VA could cause arterial dissection. The floor of cavity created after removal was carefully investigated.

Fig. 1 Preoperative cervical spine x-rays showed mild degenerative changes and collapse of the disc space at C5/6 level (Fig. 1A arrow). Although there was no clear spinal instability, neck flexion created kyphotic angle at C5/6 level, indicating hypermobility, compared with other levels (Fig. 1B). Extension x-ray revealed well-aligned C5/6 segment (Fig. 1C). A: neutral x-ray, B: flexion x-ray, C: extension x-ray.

Fig. 2 Preoperative computed tomographic angiography (CTA) of the vertebral artery (VA) (A: sagittal view, B and C: coronal view, D and E: axial view) and coronal image of MRI (T2WI) obtained with his head maintained neutral showed severe circumferential stenosis of the dominant left VA with medial displacement by an exophytic bony structure (arrow; Figs. 2A, 2B, and 2D–2G) and compression toward lateral direction by lateral disc bulging (arrowhead; Figs. 2F, and 2G) at the C5 transverse foramen. Osteophyte formation and osteoarthritis of C5/6 facet joint are showed (arrowhead; Fig. 2C), indicating spinal degenerations.

Fig. 3 Preoperative cerebral angiography with neutral neck position demonstrated severe stenosis (arrow) of the left vertebral artery (VA) at the C5/6 level (Fig. 3A) and a right VA terminating at the posterior inferior cerebellar artery (PICA) (arrowhead) (Fig. 3C). Dynamic cerebral angiography with his head rotated to the left or extended showed the left VA was completely occluded at the C5/6 level and the distal VA flow was weakly filled by collateral circulation from muscular branches (Fig. 3B). AP view: (Anterior-Posterior view).
The facet capsule was breached and articular cartilage was exposed. Judging from these intraoperative findings, the lesion appeared to be directly connected to the C5/6 facet joint. Therefore, we concluded that it had arisen from the facet joint (Fig. 4B). Spinal fusion was deemed unnecessary because of lack of significant spinal instability. Histological examination revealed a benign osteochondroma with mature bony trabeculae and bone marrow capped by cartilage tissue. Sarcomatous transformation was absent. Endochondral ossification intervened between the bone and cartilage (Figs. 5A–5C).

The patient recovered well and his neurological symptoms completely resolved immediately after surgery. CTA obtained 6 months after surgery showed no focal stenosis of the left VA and no residual or recurrent tumor (Figs. 6A–6D).

Discussion

VBI is rarely symptomatic because most patients have adequate collateral flow compensation. To be symptomatic, the contralateral VA and anterior cerebral circulation must be insufficient to provide compensatory flow. In the present case, dynamic DSA demonstrated that the left dominant VA was completely occluded at the C5 level with the leftward head rotation or extension. Distal blood flow after occlusion supplied by muscular branches was considered to be insufficient in light of the contralateral hypoplastic VA terminating in the PICA.

RVAS, also known as Bow Hunter syndrome, is VBI induced by physiological head rotation. Typically, head rotation may provoke mechanical stretching and kinking of the contralateral VA at the C1-2 level (V3 segment), which is followed by external compression of the VA by ligamentous insertions or deep fascia at the scalenovertebral angle.
Other less frequent causes include osteophytes of the uncovertebral joint, disc herniation, and spondylolisthesis at the C3-6 level where the VA passes through the transverse foramen (V2 segment).\textsuperscript{1-3,10,11} Neurological symptoms of V2 RVAS are provoked by ipsilateral head rotation or neck extension in most cases, contrary to classic RVAS at the C1/2 level. To the best of our knowledge, a spinal tumor has never been implicated as the cause of RVAS in the V2 segment. In the present case, the tumor seemed to originate from facet joint and its mobility might have led to intermittent symptoms related to head rotation or neck extension. For RVAS at the V2 segment, anterior cervical decompression with or without fusion is a reasonable approach.\textsuperscript{12,13} Resection of osteophyte or disc herniation compressing VA and unroofing of the transverse foramen are key maneuvers.

Osteochondroma is the most common benign osseous tumor and usually arises at the metaphysis of the long bone.\textsuperscript{14} It basically grows in line with bone growth and stops increasing in size with skeletal maturity. The tumor, therefore, tends to be symptomatic with an average age of around 30 years.\textsuperscript{15} In several previous reports, the appearance of symptoms at a much later age and progression of symptoms are explained by possible growth of the tumor beyond skeletal maturity,\textsuperscript{16,17} however, this abnormal tumor growth is unlikely. At the same time, we need to consider the possibility of sarcomatous transformation if we detect continuous growth of the tumor, however, this was ruled out in the present case based on histological assessment.\textsuperscript{17}

### Table 1 Summary of reported cases of vertebrobasilar insufficiency associated with spinal osteochondroma

| Authors & Year       | Age & Sex | Tumor type     | Involved spinal elements | Presenting symptoms (VA-related symptoms) | State of VA | Effect of head rotation on VA | Treatment | Surgical outcome |
|----------------------|-----------|----------------|---------------------------|------------------------------------------|-------------|-------------------------------|-----------|-----------------|
| George et al., 1989\textsuperscript{45} | 30, male  | multiple hereditary | C2 transverse process, lamina, articular facet | C2 neuralgia, tinnitus, dizziness, blurred vision (tinnitus, dizziness) | Ipsi; complete occlusion, Con; dominant and good filling to brain stem | No | Tumor removal via anterolateral approach without fusion | Complete recovery |
| Altfa et al., 2013\textsuperscript{39}    | 14, male  | multiple hereditary | C1 lateral mass          | Dizziness, headache, transient motor weakness and facial droop (all symptoms) | Ipsi; complete occlusion, Con; acute dissection | No | No surgery and antiplatelet medication for acute dissection | Complete recovery |
| Zhang et al., 2015\textsuperscript{56}   | 19, female | solitary         | C1 lateral mass          | Vertigo, nausea, vomiting, headache (all symptoms) | Ipsi; narrowing by 50%, Con; good filling to brain stem | Yes | Tumor removal, otherwise no description provided | No progression |
| Fadili et al., 2014\textsuperscript{21}  | 59, male  | solitary         | C4 posterior arch       | Coma due to bilateral cerebellar infarction and hydrocephalus (all symptoms) | Ipsi; complete occlusion, Con; dominant and intact | No | Ventriculostomy and posterior fossa decompression | Complete recovery |
| Present case         | 71, male  | solitary         | C5-6 articular facet     | Dizziness on neck rotation & extension (all symptoms) | Ipsi; complete occlusion, Con; hypoplastic terminating in the PICA | Yes | Anterior approach without fusion | Complete recovery |

Con: Contralateral, Ipsi: Ipsilateral, PICA: posterior inferior cerebellar artery, VA: vertebral artery.
from the tumor, which was occupying most of the C4 transverse foramen. Only one case reported by Zhang et al. demonstrated a dynamic change of the VA flow on neck motion during dynamic angiography like the present case. However, a clear relationship between neck movement and clinical deterioration was not revealed. The other three cases presented with complete irreversible VA occlusion. Therefore, RVAS induced by osteochondroma is first described in this report. Early exploration and detection of the tumor are essential once VBI becomes symptomatic. Presence of a tumor should be considered as a potential cause of VBI for prevention of stroke or irreversible VA occlusion. Conservative, surgical and more recently endovascular intervention have been generally proposed for VBI for prevention of stroke or irreversible VA occlusion.

The cause of VBI in the present case was a compressive osseous tumor associated with spinal degenerations and spinal mobility in the subaxial cervical spine. Complete surgical resection of the tumor and decompression of the VA without fusion was curative. The treatment modality should be selected by careful evaluation of pathology, such as the cause of compression, location, or spinal stability.

**Conclusion**

We reported a rare case of RVAS associated with spinal osteochondroma arising from the C5/6 facet joint. Imaging studies localized the lesion and the dynamic changes of the vertebrobasilar circulation were clearly revealed. Histological evaluation was necessary for the diagnosis. After surgical resection, the patient neurologically recovered well without tumor recurrence.

**Conflicts of Interest Disclosure**

The authors have no conflicts of interests.

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