Rotational vertebrobasilar insufficiency due to compression of a persistent first intersegmental vertebral artery variant: case report

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Rotational vertebrobasilar insufficiency, or bow hunter’s syndrome, is a rare cause of posterior circulation ischemia, which, following rotation of the head, results in episodic vertigo, dizziness, nystagmus, syncope, and so forth—when a patient’s head is rotated. Various case reports and case series describe the clinical entity and the role of vertebral artery decompression or cervical fusion.\(^2\),\(^5\),\(^13\) Dynamic occlusion of the vertebral artery accounts for the clinical syndrome, and the site of occlusion can seen throughout the course of the vertebral artery, with the V\(_2\) and V\(_3\) segments being the most common, respectively.\(^5\) Decompression of the artery at the site of dynamic occlusion can lead to complete resolution of symptoms in most cases.\(^5\),\(^13\)

In this report we describe a patient with RVBI secondary to a dominant persistent first intersegmental artery (PFIA), an anatomical variant of the vertebral artery at the atlantoaxial junction. In the PFIA, which is rare with a reported incidence of 0.01% in North America,\(^6\) the vertebral artery takes a low course under the arch of C-1 rather than over, bypassing the foramen transversarium of C-1.\(^6\),\(^7\),\(^12\) This is often seen in conjunction with developmental anomalies of the atlas including cranialization of the lateral mass on the side of the variant course.\(^11\)

**Case Report**

**History and Preoperative Assessment**

A 38-year-old male began experiencing spells of dizziness and near syncope upon rightward head turning. He found that the symptoms were reproducible, especially when trying to parallel park, and would resolve with neutralization of his head position. Brain MRI was initially...
performed, showing an incidental pituitary macroadenoma but no other abnormalities. Subsequent vascular imaging including CT angiography (CTA) and MR angiography (MRA) showed a hypoplastic right vertebral artery thought to be a congenital variant and a contralateral dominant left vertebral artery (Fig. 1A). The course of the left vertebral artery was consistent with a PFIA variant (Fig. 1B). The atlas was also noted to be grossly maldeveloped, with partial cranialization of the lateral masses as well as incomplete fusion of the posterior arch.

Diagnostic 4-vessel and digital subtraction angiography (DSA) with bilateral head turning was performed next. On rightward head turning only, there was a significant reduction in flow from the left vertebral artery into the posterior circulation (Fig. 1C and 1D). Dynamic imaging revealed likely compression of the PFIA as it coursed under the left-sided posterior arch of C-1. During this technique, the patient reproduced his symptoms of dizziness and near syncope, providing clinical evidence congruent with imaging findings. The diagnosis of rotational vertebrobasilar insufficiency, or bow hunter’s syndrome, was then confirmed and the patient was started on aspirin therapy.

**Operative Course**

The patient was placed prone with his head flexed, and a midline incision was made from the inion to the spinous process of C-2. Unilateral dissection of the left incomplete posterior arch of C-1 was then performed with great care. The vertebral artery was identified, and the dissection along the left incomplete arch of C-1 was carried laterally to the transverse foramen in the subperiosteal plane. Intraoperative Doppler ultrasonography was used to identify the vertebral artery, which with relative ease was found coursing below the arch as expected. Microdissection was used to confirm that no fibrous bands were present and causing soft tissue compression of the artery. The artery was compressed under a bony prominence emerging from the inferior aspect of the C-1 posterior arch. The C-1 hemilamina was then drilled with great care to the level of the C-1 transverse foramen, which the artery did not enter. Removal of this hemilamina and the associated compressive bony prominence allowed for unimpeded mobility of the vertebral artery on direct visualization and minimal manipulation. Finally, the Mayfield skull clamp was released intraoperatively, and the patient’s head was manually rotated while using Doppler ultrasonography to confirm adequate decompression.

**Postoperative Course**

Postoperative CT imaging was performed to verify the intended goals of the operation. On this study, the left hemilamina that had been seen preoperatively (Fig. 2 left) was completely removed (Fig. 2 right). This provided indirect evidence that there was good decompression of the patient’s variant vertebral artery. Immediately postoperatively, the patient reported expected rigidity and muscular spasm preventing him from performing provocative movement of his neck for the first few days. After this initial period, he was able to perform extended rightward head turning and experienced complete resolution of his preoperative symptoms.

**FIG. 1.** A: An MR angiogram demonstrating a dominant left vertebral artery (black arrow) and hypoplastic right vertebral artery (black arrowhead). B: Reconstructed CT angiogram showing PFIA (blue arrow) compared with a usual anatomical vertebral artery course (red outline). C: Formal left vertebral angiogram in a neutral position. D: Left vertebral artery on rightward head turn, revealing a significant decrease in blood flow with associated reproduction of the patient’s symptoms.

**Discussion**

Vertebrobasilar insufficiency is thought to be due to a combination of atherosclerotic disease and hypoperfusion associated with postural hypotension, exercise, or dehydration. Rotational vertebrobasilar insufficiency, or bow hunter’s syndrome, is a condition characterized by symptoms of vertebrobasilar insufficiency provoked by head turning usually in one particular direction. The association between head rotation and vertebrobasilar insufficiency was first described in 1933 by de Kley and Versteegh in their work on Ménière’s syndrome. However, it was not until 1978 that this entity was described in isolation and given its designation of “bow hunter’s stroke.” Sorensen presented a case of brainstem infarction, or modified Wallenberg syndrome, which had developed in an archer. The proposed mechanism of infarction was dynamic vertebrobasilar occlusion due to extreme or prolonged head rotation during archery sessions.

Vertebral artery anomalies at the craniocervical junction are not uncommon and occur in 0.42%–10% of the population. Permanent first intersegmental artery is one such anatomical variation of the vertebral artery thought to affect 0.01% of the US population. Most commonly, the vertebral artery arises from the subclavian arteries bilaterally and enters the foramina transversarium at the level of C-6. Upon exiting the C-1 transverse foramen, the artery traverses medially above the C-1 lamina along the
sulcus arteriosus prior to piercing the dura mater and entering the foramen magnum. With PFIA, the vertebral artery does not course through the foramen transversarium at C-1; instead, it courses medially between C-1 and C-2 and enters the canal at this level on a course through the foramen magnum. It is not usually known to cause clinical symptoms. Another variant at this level includes fenestration of the artery at C-1, in which the vertebral artery branches after leaving the foramen of C-2 and sends one branch through the foramen transversarium of C-1 on its usual course and another between the C-1 and C-2 arches. Both of these variants present a challenge in the surgical approach to C1–2 fusions as the variant arteries both traverse the placement site for lateral mass screws.

To our knowledge, this is the first report of RVBI due to a PFIA caused by compression of the artery as it courses underneath the posterior arch of C-1. We were able to witness this phenomenon in real time with DSA, which has become the gold standard in diagnosis. Surgical treatments have been described for vertebrobasilar insufficiency primarily involving decompression alone, in situ atlantoaxial fixation, or decompression combined with posterior cervical fusion. Factors determining the decision to fuse include the extent of bony decompression required, patient age, lifestyle, comorbidities, and patient preference. In patients who do not undergo arthrodesis, subsequent cervical instability and recurrent rotation-induced compression are the primary concerns. In addition, patients with RVBI commonly have a diminutive contralateral vertebral artery, and any compromise of the dominant vertebral artery during surgical decompression could have dire consequences. Endovascular vertebral artery stenting is also being investigated as a treatment strategy, though these patients must have symptomatic compression in the V₅ segment.

Posterior cervical fusion is not always described as part of the initial surgical treatment. For unilateral dynamic occlusion of the vertebral artery, decompression alone from either an anterior or posterior approach may be sufficient for symptomatic relief. However, some have described patients presenting with bilateral dynamic occlusion, in which fusion is necessary to prevent rotation since all of the posterior circulation supply is at risk in these rare cases. Fusion may also be indicated initially if decompression is extended to the point of destabilizing the atlantoaxial complex or if there is significant symptomatic spondylosis or instability on preoperative assessment. If not already available, vascular imaging studies should be obtained prior to surgical intervention if fusion is planned to guide the hardware trajectory. Computed tomography angiography is often recognized as superior to MRA as the relationship of the bony elements and vertebral arteries is clearly visualized, and it can be used for intraoperative hardware navigation.

We were able to carefully decompress the bony elements overhanging the PFIA from a posterior approach to address the source of the dynamic vascular compression. After a discussion with the patient, we decided against fusion to preserve atlantoaxial rotation given his age and active lifestyle; we were aware that any inadvertent injury to the artery during the fusion would likely be detrimental since he lacked a contralateral vertebral artery. For this patient, the surgical treatment led to a complete clinical response; he has not had any symptoms postoperatively. He has been followed up for 2 months postoperatively without complication or adverse effects from the procedure. Given his symptomatic improvement, we have not performed repeat dynamic imaging.

Conclusions

Here we describe a patient with dynamic vertebrobasilar insufficiency, or bow hunter’s syndrome, secondary to a dominant persistent first intersegmental variant of the vertebral artery. To our knowledge, this is the first description of this clinical entity being caused by this anatomical variant. Diagnosis was greatly aided by dynamic DSA and would be recommended in the workup of this condition. Decompression of the artery at C-1 at the site of dynamic impingement of C-1 was performed, and the patient showed complete resolution of his preoperative symptoms.

References


**Disclosures**
Drs. Koch and Ozturk are consultants for DePuy Synthes.

**Author Contributions**
Conception and design: Ozturk, Buch, Madsen, Vaughan, Koch. Acquisition of data: Buch, Madsen, Vaughan, Kung. Analysis and interpretation of data: all authors. Drafting the article: Ozturk, Buch, Madsen, Vaughan, Koch. Critically revising the article: Ozturk, Buch, Madsen, Vaughan, Koch. Reviewed submitted version of manuscript: all authors. Study supervision: Ozturk, Kung.

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