Delayed Infarction of Medullar and Cerebellum 3 Months after Vertebral Artery Injury with C1-2 Fracture: Case Report

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The clinical manifestations of vertebral artery (VA) injury (VAI) after cervical trauma range from asymptomatic to fatal cerebral infarction. Thrombotic occlusion and embolization to the distal arteries can cause cerebellar and brainstem infarction within days after trauma. We report a 64-year-old man who underwent arthrodesis surgery for C1 and C2 fractures. He had left VAI at the C2 transverse foramen site but was asymptomatic. The patient experiences brainstem and cerebellar infarction 3 months after injury to the VA, and we are here to discuss the treatment of VAI after cervical trauma.

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KEY WORDS: Vertebral artery · Cervical spine · Trauma · Ischemia · Clinical presentation.

Introduction

Traumatic occlusion of a vertebral artery (VA) is a rare event. Cervical fracture is highly associated with VA injury (VAI). In traumatic VAI, thrombotic occlusion and embolization to the distal arteries can cause cerebellar and brain stem infarction. Therefore, early diagnosis and treatment are necessary to decrease the risk of serious complications.

The clinical manifestations of VAI range from asymptomatic to fatal cerebral infarction. The literature also reports the occurrence of a delayed infarction a few days later.

We report a case of brain stem and cerebellar infarction 3 months after VAI and discuss about management of VAI.

Case Report

Here, we present a case of a 64-year-old man complained posterior cervical neck pain and bilateral shoulder pain after vehicle accident at April. He was ambulance driver at the time of the accident. The patient had a history of hypertension and was on medication. There was scalp laceration on vertex and no other trauma was evident. Neurological examination showed no consciousness, motor and sensory abnormality. A computed tomography (CT) scan of the cervical spine demonstrated C1 anterior, posterior arch, C2 fractures and posteriorly displaced odontoid process (Figure 1). A magnetic resonance imaging (MRI) of the cervical spine showed no cord signal change except pre-vertebral soft tissue swelling and posterior soft tissue injury at C1/2 level. A fragmented odontoid process was holding back the spinal cord. There was degenerative spondylitis in the C4–5–6–7 levels. The CT angiography scan and 3-dimensional (3D) reconstruction showed occlusion of the left distal vertebral artery in the transverse foramen at the level of C2. (Figure 2). On the following days, he underwent an arthrodesis O–C1 lateral mass-C2 pedicle - C3 pedicle fixation and on-lay graft (Figure 3). There was a...
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Fracture in the C1 posterior arch during the operation. The C1/2/3 site was very unstable due to posterior ligament injury.

The patient was discharged without any complication after surgery. Patients did not undergo preventive treatment such as aspirin for VAI. The patient complained of occipital pain for up to 3 months postoperatively, but there were no other abnormalities.

The patient re-visited with dysphasia on July 31st. The patient drank a lot of alcohol during the three days, and he said that he sweated a lot. The maximum temperature for three days was about 30°C. Neurological examination showed a clear consciousness, no motor and sensory abnormality. The patient was unable to swallow, the right pupil was 5 mm, and the left pupil was 3 mm. Uvular was deviated to the right.

Left lateral & posterior medulla, left cerebellar hemisphere

FIGURE 1. Computed tomography show fracture of C1 and C2. (A) Axial view show fracture lines (arrow) at anterior and posterior arch of atlas. (B) Sagittal reconstruction show displacement and angulation of odontoid process (arrow).

FIGURE 2. Three dimension reconstruction of vertebral artery computed tomography angiography show occlusion (arrow) at the C2 transverse process level.

FIGURE 3. Postoperative cervical lateral X-ray showed occipital-C1 lateral mass-C2 pedicle-C3 pedicle screw fixation and posterior on-lay bone graft.
FIGURE 4. Diffusion magnetic resonance imaging show restriction (arrow) at the level of medullar (A) and cerebellar (B).

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and cerebellar vermis on brain MRI with Diffusion imaging showed diffusion restriction (Figure 4). The patient underwent conservative treatment including heparin and aspirin under the diagnosis of left lateral medullary syndrome.

One month after the onset of symptoms, the patient was discharged with improved dysphagia.

The patient was followed for 40 months after the occurrence of stroke. The dysphagia has been completely restored and there is no motor and sensory abnormality besides neck pain.

Discussion

This is a case of VA area infarction that occurred 3 months after surgical treatment of C1 and 2 fractures with VAI in a 64-year-old man. Dehydration and hypercoagulability due to hot summer weather and excessive drinking was presumed to be the causative factors. Patients did not undergo preventive treatment such as aspirin for VAI. Here, we briefly review VAI after cervical trauma and discuss prophylactic aspirin.

The incidence of VAI after trauma varies widely. It is reported that about 2% of all trauma patients, 20% of head injured patients, from 13% to 46% of blunt cervical trauma, and 7% of cervical fractures are VAI. Bilateral VAI are estimated to occur at approximately 10% of the frequency of unilateral VAI. Clinical considerations are needed because posterior circulatory ischemia may result.

VAI is classified as traumatic and non-traumatic depending on the cause. Traumatic injuries are classified by penetrating injury and blunt trauma, and non-traumatic VAI occur when sudden hyperextension and rotation of the neck occurs. VAI commonly accompany blunt cervical trauma. This complication has also occurred as a result of yoga exercises, calisthenics, football injuries, bow hunting. Structurally, it occurs on fixed segment and mobile segment junctions. The C6 transverse foramen is the most common and the C1/2 site is the second most common. It is common in children with V2 segments and adults with V3 segments. In this case, the C2 transverse foramen, the border between V2 and V3, is the VAI site.

Pathologic finding in the autopsy revealed fatal dissection of the vertebral arteries. In such cases, dissection or occlusion of the vertebral or basilar arteries from thromboembolism caused brainstem ischemia and subsequent sudden death of the patient. To be noted that even small traumatic lesions in the vertebral arteries could cause brain infarction in the absence of occlusion of the vertebral or basilar arteries; subsequent death would occur from complications.

Clinical features range from asymptomatic to fatal posterior circulation infarction. Unilateral VA occlusion is rarely symptomatic because of collateral blood supply from the contralateral VA and the circle of Willis. Moreover, injuries to the non-dominant VA are not likely to be symptomatic. However, unlike unilateral VA occlusion, in which only 20% of patients are symptomatic, most patients with bilateral VA occlusion are symptomatic. Bilateral or dominant VA occlusion may cause rapid and fatal ischemic damage to the cerebellum and brain stem.

Symptoms include ischemia-induced symptoms in the brain stem, cerebellum, and occipital lobe. It can cause mild headache, sensory abnormality, muscle weakness, dizziness, disappearance of co-ordination, visual defect, Horner syndrome and lateral medullary syndrome, loss of con-
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There is a controversy about aggressive screening in VAI because it is relatively small compared to the number of head trauma patients. VAI after blunt cervical trauma was previously considered to be rare. It is also argued that the early diagnosis of VAI due to aggressive screening does not significantly affect the outcome of the treatment.13 The recent increase in incidence is mainly due to improvements in diagnosis, particular in imaging techniques such as digital subtraction angiography (DSA), MR angiography (MRA), and CT angiography (CTA). CTA in particular is a first-choice diagnostic tool for VAI because it is non-invasive with respect to the vascular system, is easy to use, and has a high detection rate.4 In this reason, CTA is a recommended for patients who are eligible for modified Denver Screening Criteria for suspected VAIs (Table 1).5

Treatment of VAI includes anticoagulation, antiplatelet, endovascular intervention, surgical vessel sacrificing, and no treatment depending on the situation. However, there are no clear guidelines yet. According to several authors, recurrent transient ischemic attacks, stroke or death could occur within weeks unless anticoagulation or antiplatelet use.1,2,4,5,13 Desouza et al.4 recommend that patients with VAI start anticoagulation and then change to antiplatelet, regardless of the presence or absence of VAI. However, Harrigan et al.5 have recommended a multicenter, randomized, and prospective study of anticoagulation and antiplatelet efficacy for asymptomatic patients.

In this case, asymptomatic patient developed posterior circulation ischemia 4 months after VAI. The authors recommend antiplatelet because the use of antiplatelet will be more beneficial given the age of 64 and the risk factors for stroke, such as drinking, hypertension, and smoking.

**Conclusion**

This is a case of VA area infarction that occurred 3 months after surgical treatment of C1 and 2 fractures with VAI in a 64-year-old man. Patients with asymptomatic VAI with risk factors may benefit from antiplatelet treatment rather than no treatment.

*The authors have no financial conflicts of interest.*

**REFERENCES**

1) Beletsky V, Nadareishvili Z, Lynch J, Shuaib A, Woolfenden A, Norris JW. Cervical arterial dissection: time for a therapeutic trial? *Stroke* 34:2856-2860, 2003
2) Biffl WL, Moore EE, Elliott JP, Ray C, Offner PJ, Franciose RJ, et al. The devastating potential of blunt vertebral arterial injuries. *Ann Surg* 231:672-681, 2000
3) Chung D, Sung JK, Cho DC, Kang DH. Vertebral artery injury in destabilized midcervical spine trauma; predisposing factors and proposed mechanism. *Acta Neurochir (Wien)* 154:2091-2098, 2012
4) Desouza RM, Crocker MJ, Haliasos N, Rennie A, Saxena A. Blunt traumatic vertebral artery injury: a clinical review. *Eur Spine J* 20:1405-1416, 2011
5) Harrigan MR, Hadley MN, Dhall SS, Walters BC, Ararabi B, Gelb DE, et al. Management of vertebral artery injuries following non-penetrating cervical trauma. *Neurosurgery* 72 Suppl 2:234-243, 2013
6) Iwase H, Kobayashi M, Kurata A, Inoue S. Clinically unidentified dissection of vertebral artery as a cause of cerebellar infarction. *Stroke* 32:1422-1424, 2001
7) Jang D, Kim C, Lee SJ, Kim J. Delayed Brain Infarction due to Bilateral Vertebral Artery Occlusion Which Occurred 5 Days after Cervical Trauma. *J Korean Neurosurg Soc* 56:141-145, 2014

**TABLE 1. The Denver screening criteria for blunt cerebrovascular injury and Denver radiological grading scale of blunt cerebrovascular injury**

| Denver screening criteria                    | Denver radiological grading scale |
|---------------------------------------------|-----------------------------------|
| Any cervical spine fracture                 | Grade I                           |
| Unexplained neurological deficit incongruent with imaging | Irregularity of vessel wall or dissection/intramural hematoma with <25% stenosis |
| Basilar cranial fracture into carotid canal | Grade II                          |
| Le Fort II or III fracture                  | Intramural thrombus or raised intimal flap or dissection/intramural hematoma with >25% stenosis |
| Cervical hematoma                           | Grade III                         |
| Horner syndrome                             | Pseudoaneurysm                     |
| Cervical bruit                              | Grade IV                          |
| Ischemic stroke                             | Vessel occlusion                   |
| Head injury with Glasgow Coma Scale score < 6 | Grade V                           |
| Haring with anoxic injury                   | Vessel transection                 |

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8) Jang JW, Lee JK, Hur H, Seo BR, Lee JH, Kim SH. Vertebral artery injury after cervical spine trauma: A prospective study using computed tomographic angiography. Surg Neurol Int 2:39, 2011
9) Medhkour A, Chan M. An unusually favorable outcome of bilateral vertebral arterial dissections: case report and review of the literature. J Trauma 58:1285-1289, 2005
10) Mimata Y, Murakami H, Sato K, Suzuki Y. Bilateral cerebellar and brain stem infarction resulting from vertebral artery injury following cervical trauma without radiographic damage of the spinal column: a case report. Skeletal Radiol 43:99-105, 2014
11) Mueller CA, Peters I, Podlogar M, Kovacs A, Urbach H, Schaller K, et al. Vertebral artery injuries following cervical spine trauma: a prospective observational study. Eur Spine J 20:2202-2209, 2011
12) Pötsch L., Bohl J. Traumatic lesion of the extracranial vertebral artery—a note-worthy potentially lethal injury. Int J Legal Med 107: 99-107, 1994
13) Stein DM, Boswell S, Sliker CW, Lui FY, Scalea TM. Blunt cerebrovascular injuries: does treatment always matter? J Trauma 66: 132-143, 2009
14) Yoshihara H, Vanderheiden TF, Harasaki Y, Beauchamp KM, Stahel PF. Fatal outcome after brain stem infarction related to bilateral vertebral artery occlusion - case report of a detrimental complication of cervical spine trauma. Patient Saf Surg 5:18, 2011