Traumatic Basilar Artery Entrapment without Longitudinal Clivus Fracture: A Case Report and Review of the Literature

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Abstract

In blunt cerebrovascular injury, reported traumatic basilar artery occlusions have involved dissection of the basilar artery, distal embolization due to traumatic vertebral artery dissection, or entrapment of the basilar artery into the clivus fracture. To date, however, there are no reports of traumatic basilar artery entrapment without a clivus fracture. Here, we report the first case of traumatic basilar artery occlusion caused by entrapment into an originally existing bone defect. A 67-year-old man with a history of treatment for intracranial aneurysm suffered multiple traumatic injuries in a fall. On arrival at our hospital, he presented with neurogenic shock with quadriplegia. Computed tomography (CT) showed small epidural hematoma, C4-6 cervical spinous process fracture, and Th2-3 vertebral body fracture. CT angiography revealed occlusion of the basilar artery trunk. As vertebrobasilar artery dissections and clivus fracture were not observed; however, we could not elucidate the pathology of the basilar artery occlusion. On day 4, after surgery for the cervical and thoracic lesions, he exhibited consciousness disturbance. Diffusion-weighted imaging on day 5 showed hyperintensities in the brainstem and cerebellum. Basi-parallel anatomic scanning magnetic resonance imaging showed that the basilar artery, while lacking vascular wall injuries, was tethered into the clivus. Antithrombotic therapy was performed, but the patient progressed to a locked-in state. Previous head CT before the trauma revealed a bone defect already present in the clivus. We speculated basilar artery entrapment into this preexisting bone defect. We must look for basilar artery injury in trauma patients even in the absence of clivus fracture.

Key words: traumatic basilar artery entrapment, brainstem infarction, spheno-occipital synchondrosis, basi-parallel anatomical scanning

Introduction

Blunt cerebrovascular injuries (BCVIs) affecting the carotid or vertebral artery can lead to abnormal arterial narrowing, pseudoaneurysm, and occlusion, and typically occur in the setting of either generalized multiple trauma or direct craniocervical trauma. Although rare, there are some reports of traumatic basilar artery occlusion causing lethal infarction of the brainstem and severe disability or death.1–10 Reported mechanisms of traumatic basilar artery occlusion have involved dissection of the basilar artery originating from the trauma itself,11 distal embolization due to traumatic vertebral artery dissection,12 or entrapment of the basilar artery into a longitudinal clivus fracture.1–10,13–16 In the absence of clivus fracture, however, to the best of our knowledge, there are no reports of traumatic basilar artery entrapment. Here, we report the first case of traumatic basilar artery occlusion caused by entrapment into an originally existing bone cortex defect at the spheno-occipital synchondrosis, and review the relevant published literature.
Case Report

A 67-year-old man fell to the ground from the cargo bed of a truck (height of approximately 1 m), and was initially transferred to another emergency hospital. He suffered from neurogenic shock with quadriplegia, and was treated with an adrenergic vasopressor. He was transferred to our hospital for multidisciplinary treatment. On arrival at our hospital, the patient was alert with a Glasgow Coma Scale score of 14 (E3V5M6). His vital signs were respiratory rate 23 breaths/min, blood pressure 106/69 mmHg (dopamine, 5.5 μg/kg/min), and heart rate 70 bpm (sinus rhythm). His oxygenation on pulse oximetry (SpO₂) was 99% (O2 1L). Computed tomography (CT) of the head revealed a small epidural hematoma at the frontal bone of the skull. He had a past history of bifrontal craniotomy for surgical clipping of an intracranial aneurysm 3 years earlier at another hospital. He also exhibited a C4-6 cervical spinous process fracture, Th2-3 vertebral-body fracture, bilateral first rib fracture, and sternal fracture. T2-weighted magnetic resonance imaging (MRI) showed high-intensity areas at the C3-4 and Th2 spinal cord. Three-dimensional CT angiography (3D-CTA) as part of a whole-body CT scan revealed occlusion of the basilar artery trunk and poor description of the left vertebral artery distal to the posterior inferior cerebellar artery and the right vertebral artery proximal to the bifurcation. The lower half of the basilar artery trunk was entrapped in a minor bone defect in the clivus (Fig. 1A, C–E). In contrast, no abnormal findings, such as arterial dissection were observed in the vertebral artery near the cervical vertebral fracture. We could not recognize any obvious clivus fracture on multi-planar CT images. The patient presented no neurological abnormality except for quadriplegia. Because of ongoing management, including surgery for multiple traumas, antithrombotic therapy was not indicated. On day 3, the patient underwent posterior fusion surgery for the cervical and thoracic lesions, and went into hypovolemic shock due to intraoperative bleeding. His systolic blood pressure decreased to <80 mmHg several times. After the surgery, he exhibited delayed awakening from anesthesia with severe consciousness disturbance. On day 4, head CT showed no evidence of intracranial hemorrhage but did show a small low-density area on the pons and cerebellum. On day 5, diffusion-weighted imaging showed hyperintensities in the area of the pontine arteries and the left anterior inferior cerebellar artery (Fig. 2A, B). We retrospectively recognized that the lower half of the basilar artery trunk was entrapped in a minor bone defect in the clivus (Fig. 2C, D).

Fig. 1 Computed tomography (CT) images of the head at admission showed occlusion of the basilar artery trunk. (A) Poor description of the left vertebral artery distal to the posterior inferior cerebellar artery (C), and the right vertebral artery proximal to the bifurcation (E). The lower half of the basilar artery trunk was entrapped in a minor hole in the middle part of the clivus (B, D).

Fig. 2 Magnetic resonance imaging (MRI) of the head on day 5 (A, B) showed hyperintensities in the area of the pontine arteries and the left anterior inferior cerebellar artery. Basi-parallel anatomic scanning MRI (BPAS-MRI) on day 6 (C) revealed that the outer contour of the basilar artery was narrowed at the middle portion of the clivus (black arrow). BPAS-MRI on day 18 (D) showed that the basilar artery trunk was tethered to the clivus (white dotted circle).
middle part of the clivus, which was visible on the 3D-CTA performed at admission (Figs. 1B and 3B). We confirmed stabilization of the hemorrhagic lesions at other trauma sites and started antithrombotic therapy with heparin and aspirin immediately after the diagnosis. Basi-parallel anatomic scanning MRI (BPAS-MRI) findings on day 6 revealed that the outer contour of the basilar artery narrowed at the middle portion of the clivus (Fig. 2C). On day 10, 3D digital subtraction angiography (DSA) confirmed retrograde blood flow from the right posterior communicating artery to points distal to the basilar artery occlusion. Sagittal BPAS-MRI images on day 18 revealed that the basilar artery trunk with a normal outer contour was tethered to the middle portion of the clivus (Fig. 2D). The patient progressed to a locked-in state, and was transferred to another hospital for rehabilitation on day 19. Previous head CT performed at another hospital at the time of aneurysmal clipping 3 years prior to this injury revealed that the bone defect in the middle clivus was present at that time (Fig. 3A). The location of the bone defect was highly suggestive of a remnant of sphenoid-occipital synchondrosis.

Discussion

The course of our case suggested two important clinical issues. First, it should be noted that the traumatic basilar artery occlusion occurred without a clivus fracture. Second, the combination of radiological modalities was useful in the differential diagnosis of basilar artery occlusion.

About 15 previous cases of traumatic basilar artery entrapment can be retrieved by searching the PubMed electronic database (Table 1). In our review, 14 involved a longitudinal clivus fracture, and the remaining case involved a nondisplaced clivus fracture. The typical mechanism of traumatic basilar artery entrapment is understood as follows: the basilar artery is pressed against the fissure of a longitudinal clivus fracture, the fracture is closed, and the basilar artery is incarcerated. This injury mechanism is thought to explain why traumatic basilar artery entrapment cannot usually occur without a longitudinal clivus fracture. Our case, in contrast, exhibited no clivus fracture, but a minor bone defect in the clivus was observed. We suspected that this bone defect had occurred as a remnant of sphenoid-occipital synchondrosis in the process of endochondral ossification. After birth, the clivus develops from a sphenoid bone at the top and an occipital bone at the bottom. Between these bones is the spheno-occipital synchondrosis. It has been reported that complete closure of this synchondrosis in males occurs between 13 and 25 years of age. However, complete holes in this location have also been reported in several cases associated with spontaneous cerebrospinal fluid rhinorrhea. The spheno-occipital synchondrosis develops through endochondral ossification without fusion points, which could explain the occasional development of bone defects and eventual dehiscence at this site. In our case, moreover, CT findings of the cervical vertebral fracture indicated a cervical extension injury pattern. It was suggested that the spinal cord had been towed downward. From these
findings, the following mechanisms were presumed to have caused this case. A laceration occurred in the dura due to external force from the fall. The brainstem was towed downward by the traction of the spinal cord. The basilar artery was compressed between the brainstem and the clivus bone, and was coincidentally entrapped in the bone defect of the clivus. Additionally, we speculated that the entrapment of the basilar trunk has progressively led to a retrograde thrombosis or dissecting occlusion and delayed pontine perforators infarction (Fig. 3C).

The combination of radiological modalities was useful in the differential diagnosis of basilar artery occlusion. Because CTA is affected by blood flow and the contrast agent is visualized in the vascular inner luminal regions, it is difficult to visualize the vessels themselves. Evaluating the outer contour of the basilar artery by BPAS-MRI enables us to distinguish whether pathology is due to arterial dissection. Traumatic basilar artery occlusion due to entrapment is often diagnosed based on radiological findings showing a longitudinal clivus fracture and obvious invagination of the basilar artery. In our case, although CTA and MRA revealed the occlusion of the basilar artery, bone scan imaging of the cranial base did not show the clivus fracture. There were no findings suggesting the dissection of vertebrobasilar arteries such as intimal flaps, double lumen, or the pearl and string sign. BPAS-MRI images of this case revealed that the basilar artery was tethered into the clivus in the absence of any vascular wall injury or narrowing. These findings supported our speculation of basilar artery entrapment in an originally existing bone defect. In similar cases in the future, further evaluation for the pathology of basilar artery occlusion may be

Table 1  Summary of patients with traumatic basilar artery entrapment

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Age/sex</th>
<th>GCS score</th>
<th>Diagnostic method</th>
<th>Clivus fracture</th>
<th>Basilar artery</th>
<th>Treatment</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fang (2012)</td>
<td>39 M</td>
<td>3 (Drunk)</td>
<td>CT, CTA</td>
<td>Longitudinal</td>
<td>Severe stenosis → Occlusion</td>
<td>AT</td>
<td>Completely recovered</td>
</tr>
<tr>
<td>Garcia-Garcia (2012)</td>
<td>37 M</td>
<td>15</td>
<td>CT, CTA</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>AT</td>
<td>Mild left hemiparesis</td>
</tr>
<tr>
<td>Bala (2004)</td>
<td>46 M</td>
<td>15</td>
<td>CT, CTA, MRA</td>
<td>Longitudinal</td>
<td>Severe stenosis</td>
<td>AT</td>
<td>Mild left hemiparesis</td>
</tr>
<tr>
<td>Khanna (2010)</td>
<td>55 M</td>
<td>8 (Sedated)</td>
<td>CT, CTA</td>
<td>Longitudinal</td>
<td>Severe stenosis</td>
<td>Not described</td>
<td>Left hemiparesis</td>
</tr>
<tr>
<td>Taguchi (2000)</td>
<td>52 M</td>
<td>3 (Drunk)</td>
<td>CT, MRA</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>AT</td>
<td>Quadriplegia</td>
</tr>
<tr>
<td>Present case (2017)</td>
<td>67 M</td>
<td>3</td>
<td>CT, CTA, BPAS-MRI</td>
<td>None</td>
<td>Occlusion</td>
<td>AT</td>
<td>Locked-in state</td>
</tr>
<tr>
<td>Wang (2017)</td>
<td>59 M</td>
<td>15</td>
<td>CT, CTA</td>
<td>Longitudinal</td>
<td>Focal stenosis</td>
<td>AT</td>
<td>Locked-in state</td>
</tr>
<tr>
<td>Sen-Gupta (2012)</td>
<td>67 M</td>
<td>11</td>
<td>CT, CTA</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Not described</td>
<td>Locked-in state</td>
</tr>
<tr>
<td>Kaakaji (2004)</td>
<td>50 M</td>
<td>6</td>
<td>CT, MRA</td>
<td>Nondisplaced</td>
<td>Severe stenosis</td>
<td>EVD</td>
<td>Locked-in state</td>
</tr>
<tr>
<td>Guha (1989)</td>
<td>27 M</td>
<td>3</td>
<td>CT, DSA</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Not described</td>
<td>Vascular state</td>
</tr>
<tr>
<td>Kliesch (2017)</td>
<td>Adult</td>
<td>11</td>
<td>CT, CTA, 3D-FPA</td>
<td>Longitudinal</td>
<td>Severe stenosis</td>
<td>DC</td>
<td>Death</td>
</tr>
<tr>
<td>Sato (2001)</td>
<td>56 M</td>
<td>5</td>
<td>CT, DSA</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Not described</td>
<td>Death</td>
</tr>
<tr>
<td>Anthony (1987)</td>
<td>70 M</td>
<td>3</td>
<td>Autopsy</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Not described</td>
<td>Death</td>
</tr>
<tr>
<td>Shaw (1972)</td>
<td>59 M</td>
<td>3</td>
<td>Autopsy</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Supportive care</td>
<td>Death</td>
</tr>
<tr>
<td>Sights (1968)</td>
<td>23 M</td>
<td>3</td>
<td>Autopsy</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Supportive care</td>
<td>Death</td>
</tr>
<tr>
<td>Loop (1964)</td>
<td>59 M</td>
<td>3</td>
<td>Autopsy</td>
<td>Longitudinal</td>
<td>Occlusion</td>
<td>Supportive care</td>
<td>Death</td>
</tr>
</tbody>
</table>

important to determine whether reperfusion therapy is warranted during the acute phase. A previous case report suggested that endovascular treatment might be effective for distal embolization of the basilar artery due to traumatic vertebral artery dissection.

For early diagnosis and treatment of BCVI, appropriate evaluation of the head and neck vessels based on signs or symptoms and risk factors must be performed as soon as possible. Antithrombotic therapy has been reported as the standard treatment for traumatic basilar artery entrapment. In our review, treatment has been described in nine cases and five cases including our case received antithrombotic therapy. Three cases with good neurological outcome (completely recovered or mild left hemiparesis) were diagnosed as basilar artery entrapment on the first day and started treatment. In previous case reports, bifrontal decompressive craniectomy or external ventricular drainage has been performed to control elevated intracranial pressure caused by the infarction. There are no reports, however, of direct surgical or endovascular treatment for basilar artery entrapment. In our case, we were able to detect traumatic basilar artery occlusion on cranial CTA at admission. Antithrombotic therapy was not started; however, we did not observe apparent clinical signs of basilar artery occlusion and peri-operative management. Also, we did not perform an assessment of cerebral blood flow besides CTA at admission. It was reported that most patients with BCVIs exhibit "a latent period (10–72 h)" between their injury and the onset of stroke. In our review, four cases including our case were alert at admission (GCS 14 or 15). Two cases, however, developed consciousness disturbance and proceeded to a locked-in state. Even in the absence of clinical signs, if there are equivocal findings, early evaluation of cerebral blood flow such as DSA is needed. Furthermore, immediate antithrombotic therapy for basilar artery occlusion is crucial and should be stopped only during the peri-operative period. We also speculated that the patient’s hypovolemic shock during surgery might have caused delayed cerebral infarction. In the case of multiple traumas, it is often difficult to introduce antithrombotic therapy in the acute phase. Strict blood pressure management should be considered to prevent hypovolemic, obstructive, or neurogenic shock.

**Conclusions**

We experienced a rare case of traumatic basilar artery entrapment that caused severe disability due to basilar artery occlusion in the absence of a clivus longitudinal fracture. The combination of radiological modalities was useful to detect the cause of traumatic basilar artery occlusion. Early indication of antithrombotic therapy and systemic management are essential for preventing severe cerebral infarction.

**Conflicts of Interest Disclosure**

No authors have conflicts of interest.

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