Successful Treatment of Sudden Bilateral Sensorineural Hearing Loss due to Atherosclerotic Vertebral Artery Occlusion: A Case Report

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Objective: We report a case of sudden bilateral sensorineural hearing loss caused by atherosclerotic occlusion of vertebral artery which improved after angioplasty with stenting.

Case Presentation: The patient was a 71-year-old male. He was referred to our hospital from an otolaryngologist with severe, rapidly progressing hearing loss and vertigo. Cerebral angiography revealed occlusion of the predominant left vertebral artery, suggesting ischemia in the bilateral anterior inferior cerebellar artery (AICA) territories. Urgent balloon angioplasty was performed but after additional angioplasty, acute vessel reocclusion occurred associated with intimal dissection. Rescue stenting was performed and led to recanalization. After treatment, his hearing disturbance rapidly improved.

Conclusion: Cochlear disorder caused by ischemia in the bilateral AICA regions may induce severe acute bilateral hearing loss. Acute revascularization is useful, but it is necessary to prepare stents for angioplasty and carefully evaluate whether treatment is indicated.

Keywords ▶ sudden hearing loss, anterior inferior cerebellar artery, angioplasty and stenting, intracranial atherosclerosis, internal auditory artery

Introduction

Acute sensorineural hearing impairment with vertigo is related to vestibular dysfunction, such as Meniere’s disease, vestibular neuritis, and internal otitis, in many cases. However, ischemia in the anterior inferior cerebellar artery (AICA) territory may cause vertigo or hearing impairment.\(^1\) In this study, we report a patient in whom ischemia in the bilateral AICA territories associated with atherosclerotic occlusion of the vertebral artery induced vertigo and bilateral sensorineural hearing loss with rapid progression, and acute revascularization improved the symptoms.

Case Presentation

Case: A 71-year-old male.

Complaints: Bilateral hearing impairment, vertigo, and gait disorder.

Present illness: During the daytime, vertigo was noted while working. At night, it became exacerbated, and he consulted an emergency hospital. Under a tentative diagnosis of aural vertigo, he was referred to the Department of Otorhinolaryngology. An oral agent was administered, and follow-up was performed. However, the following day, bilateral hearing impairment appeared and rapidly progressed. In addition, gait disorder was noted, and he consulted the Department of Otorhinolaryngology of another hospital. Brain MRI revealed cerebral infarction, and he was referred to our hospital.

Medical history: Hypertension and diabetes mellitus. Detailed examination of the head indicated stenosis of the left vertebral artery 5 months prior to presentation. Clopidogrel at 75 mg/day was administered.
Sudden Hearing Loss due to Vertebral Artery Occlusion

Family history: Not contributory.
Lifestyle: Smoking (20 cigarettes/day, 45 years).
Physical examination on admission: The Japan Coma Scale (JCS) score was 1 (severe bilateral hearing impairment required communication by writing). There was no facial sensory, movement, or ocular movement disorder. Horizontal rotatory nystagmus to the left and atactic gait was observed, and the results of left/right finger to nose tests were poor.
Pure-tone audiogram (PTA): Severe hearing impairment was noted (right: 92.5 dB, left: 72.5 dB [4-minute method]) (Fig. 1A).

Neuroradiological findings
On brain MRI, high-signal intensity in the bilateral middle cerebellar peduncles was noted on diffusion-weighted images (DWIs) (Fig. 2A). A similar finding was also observed on T2-weighted images (Fig. 2B). Brain MRA 5 months before onset revealed stenosis of the left vertebral artery (Fig. 3A), and occlusion of the left vertebral artery and reduction in the visualization of the basilar artery were noted on MRA on admission (Fig. 3B). Cerebral angiography revealed occlusion of the predominant-side left vertebral artery (Fig. 4A and 4B) and hypoplasia.
of the right vertebral artery peripheral to the posterior inferior cerebellar artery; the proximal basilar artery, contralateral left posterior inferior cerebellar artery, and bilateral anterior inferior cerebellar arteries were visualized (Fig. 4D and 4E). On bilateral common carotid angiography, the hypoplastic posterior communicating artery to basilar artery end was visualized (Fig. 4C and 4F). Based on these findings, occlusion of an atherosclerotic lesion of the left vertebral artery may have caused the marked ischemia in the bilateral AICA territories. As hearing had rapidly progressed, endovascular treatment was performed.

Endovascular treatment
Prior to treatment, two antiplatelet agents, clopidogrel at 300 mg and aspirin at 200 mg (loading dose), were administered. Systemic heparinization was performed to maintain
the activated clotting time (ACT) at ≥250. Under local anesthesia, a 6 Fr FUBUKI guiding catheter (Asahi Intecc Co., Ltd., Aichi, Japan) was inserted into the left vertebral artery through the right femoral artery. A 200-cm 0.014-inch CHIKAI microguidewire (Asahi Intecc Co., Ltd.) with a J-shaped tip was guided to the site of occlusion together with an Excelsior SL-10 microcatheter (Boston Scientific, Natick, MA, USA). At the site of occlusion, the microguidewire was inserted into the occluded portion and advanced up to the distal of the lesion. The microcatheter was then navigated over the wire across the occluded segment. Microcatheter angiography was performed to confirm the presence of the microcatheter in the true lumen, the microguidewire was advanced into the left posterior cerebral artery, and prolonged using an ASAHI EXTENSION NV extension wire (Asahi Intecc Co., Ltd.) to replace the microcatheter with a balloon catheter. A Gateway balloon for percutaneous angioplasty 1.5 mm in diameter and 9 mm in length (Stryker, Kalamazoo, MI, USA) was introduced over the microguidewire, positioned to the occluded lesion and inflated to a nominal pressure of 6 atm through slow inflation (1 atm/20 seconds). Recanalization was obtained, but there was still significant residual stenosis (Fig. 5A and 5D). So the balloon was changed to a Gateway balloon 2.0 mm in diameter and 15 mm in length. This balloon was guided to the stenotic site, and inflation was performed, as described above, leading to acute reclosure (Fig. 5B and 5E). Under a diagnosis of acute occlusion associated with intimal dissection, a Wingspan 2.5 mm in diameter and 15 mm in length (Stryker) was deployed/placed across the lesion as rescue treatment. Although there was still moderate degree of residual stenosis, circulation was dramatically improved. The absence of changes was confirmed by repeat angiography (Fig. 5C and 5F) after waiting for 15 minutes with the microwire in place, and the procedure was completed.
Course after treatment
MRI (DWI) (Fig. 6A) the day after treatment did not reveal any new lesions. Furthermore, MRA confirmed favorable basilar artery blood flow (Fig. 6B). After treatment, his hearing rapidly improved. PTAs (Fig. 1B) indicated improvement of hearing (right: 20 dB [4-minute method], left: 22.5 dB [4-minute method]), and atactic gait gradually improved. Dual-antiplatelet therapy with clopidogrel at 75 mg/day and aspirin at 100 mg/day was continued. In addition to the medical management of hypertension and diabetes mellitus, lifestyle management, such as smoking cessation, was initiated. There has been no recurrence. Angiography 12 months after treatment did not demonstrate restenosis (Fig. 6C and 6D).

Discussion
Acute sensorineural hearing loss is considered to be an otorhinolaryngological symptom related to idiopathic hearing impairment or Meniere’s disease. Many patients often consult the Department of Otorhinolaryngology and receive treatment. However, a similar symptom is sometimes induced by ischemic cerebrovascular disease involving the vertebrobasilar artery territory. The incidence of hearing impairment related to cerebral infarction involving the vertebrobasilar artery system ranges from 1.4% to 8%. Although it differs among studies, this disorder is not so rare. According to one previous study, infarction involving the AICA region accounted for 5.5% of cerebral infarction involving the vertebrobasilar artery system, but it accounted for 83.3% of vertebrobasilar-artery-system cerebral infarction with acute hearing loss, and posterior inferior cerebellar artery infarction accounted for 11.9%. In particular, ischemia in the AICA territories considered to be a primary etiological factor for hearing impairment.

The AICA originates from the 1/3 caudal-side area of the basilar artery, and sometimes from the 1/3 middle area or the confluence of the bilateral vertebral arteries. It courses through the lateral part of the pons, crosses near the facial and vestibulocochlear nerve at the cerebellopontine angle, and bifurcates into the lateral and medial branches. At this segment, the internal auditory artery arises from the lateral branch. The internal auditory artery, which supply the cochlea, branches from the meatal loop of the AICA, but rarely from the posterior inferior cerebellar or basilar arteries. The internal auditory artery consists of one to several terminal arteries with little collateral circulations. Ischemia here may cause irreversible cochlear disorder. On the other hand, the cochlear nerve has an abundant collateral circulation, differing from the cochlea, and may resist ischemia.

Lee et al. found that sudden hearing impairment was an important sign of AICA infarction, and concluded that hearing impairment and vertigo were caused by ischemic cochlear disorder in many patients. In the present case, thrombotic occlusion of the predominant-side left vertebral artery led to hypoplastic distal right vertebral artery-mediated blood flow involving the proximal basilar artery, contralateral left posterior inferior cerebellar artery, and bilateral anterior inferior cerebellar arteries. In addition, a retrograde collateral pathway mediated by the posterior communicating artery was not sufficient, resulting in hypo-perfusion in the bilateral AICA territories. As a result, insufficient blood flow of the internal auditory artery as a terminal artery may have caused the cochlear dysfunction, leading to bilateral hearing loss.

Lee et al. reported that hearing recovered to some degree in approximately 80% of patients with hearing impairment.
related to cerebral infarction involving the vertebrobasilar artery region, whereas it was unfavorable in severe-status patients.\(^1\) One previous Japanese study examined hearing on initial consultation and its improvement rating in 46 patients with AICA syndrome related to stenosis/occlusion of the AICA, and reported that there were no hearing recovery in 16 (44.4%) of 36 patients in whom it was possible to evaluate the improvement rating of hearing. Excluding 10 in whom it was unclear, hearing recovery was not favorable.\(^8\) In addition, an experiment using guinea pigs demonstrated that hair cells were particularly damaged due to the effects of internal auditory artery occlusion on the cochlea; internal hair cells were damaged 3 hours after occlusion, and external hair cells 4 hours after occlusion. The disappearance of organum spirale hair cells was observed 6 hours after occlusion.\(^9\) Therefore, it may be necessary to improve blood flow as early as possible in patients with severe hearing impairment.

Currently, when performing endovascular treatments for acute ischemic stroke due to intracranial large artery occlusion, mechanical thrombectomy with a stent retriever has been recommended as the first-line treatment. However, intracranial atherosclerotic stenosis (ICAS) is involved in the pathogenesis in some patients. Yoon et al. reported that ICAS was present in 22.9% of patients with acute cerebral ischemia who underwent endovascular treatment, and indicated the usefulness and safety of mechanical thrombectomy with a stent retriever as first-line treatment, followed by angioplasty with or without stenting for residual stenotic lesions.\(^10\) In the present case, the presence of an atherosclerotic stenotic lesion on the left vertebral artery had been confirmed on previous MRA, and balloon angioplasty was primary performed. For angioplasty with a balloon, slow inflation is essential, but dissection may occur in some cases; therefore, a backup stent is necessary. In Japan, the self-expandable stent Wingspan is the only type approved for ICAS. However, this device is not always available. Thus, at our hospital, stents for stent-assisted coil embolization and balloon-mounted coronary stents are always prepared through approval by the Ethics Review Board to establish a backup system although these stents have not been approved for ICAS. In Japan, it is recommended that a Wingspan stent be indicated for vascular dissection during balloon angioplasty, acute/impending occlusion as a rescue treatment option, or additional treatment after angioplasty in cases in which there is no other effective treatment method.\(^11\) In the present case, vascular dissection-related acute occlusion occurred after balloon angioplasty, requiring the use of a Wingspan stent for rescue stenting. Furthermore, ICAS-associated hemodynamic cerebral ischemia is considered to be a better indication for intracranial stents in comparison with branch atheromatous disease or thromboembolic occlusion.\(^12\) In the present case, the recovery from hemodynamic cerebral ischemia may also have led to the rapid improvement in hearing. On the other hand, atherosclerotic debris being displaced or “snow-plowed” may occur during angioplasty or stenting, inducing occlusion of perforating arteries originating from an area adjacent to a stenotic lesion,\(^13\) as indicated by the SAMMPRIS study.\(^14\) In addition, the dura around the penetrating area of the V4 segment of the vertebral artery is much thicker than in other areas, and it forms a fibrous dural band;\(^15\) therefore, stent deformity associated with head-movement-related mechanical stimuli and subsequent intimal injury-related restenosis/occlusion may occur. As such, endovascular treatment should be performed only when revascularization is considered to be more effective than best medical treatment.

### Conclusion

We describe a case of sudden bilateral sensorineural hearing loss caused by atherosclerotic occlusion of vertebral artery successfully treated by angioplasty with stenting. Insufficient blood flow of the internal auditory artery as a terminal artery may have caused the cochlear dysfunction, leading to bilateral hearing loss. For angioplasty, it is necessary to prepare stents such as a Wingspan stent. Furthermore, it is important to carefully evaluate whether treatment is indicated.

### Disclosure Statement

There is no conflict of interest regarding this article.

### References


