Subarachnoid Hemorrhage from Vertebral Arteriovenous Fistula without Perimedullary Drainage: Rare Stroke Hemorrhagic Event in a Patient of Neurofibromatosis Type 1

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Abstract

Vertebral arteriovenous fistula (VAVF), which can cause subarachnoid hemorrhage (SAH) when having a perimedullary drainage, has been reported as a rare vascular abnormality in patients with neurofibromatosis type 1 (NF-1). In addition, extracranial vertebral aneurysm (EVAn) coexisting with VAVF and NF-1 is considered rare, and further complication with SAH is extremely rare in patients. There is only one reported case of NF-1 complicated with SAH from VAVF with an EVAn. Here, we present a case of a middle-aged patient with NF-1. The VAVF accompanied by an EVAn was detected with an episode of SAH. The VAVF with an EVAn in our case was accompanied with an epidural varix, lacking of perimedullary drainage, which could be a cause for SAH. We speculate the mechanism of SAH from the VAVF with an EVAn lacking of perimedurally drainage, focusing on hemodynamic stress of the VAVF and the tissue fragility related to NF-1.

Key words: vertebral arteriovenous fistula, vertebral aneurysm, neurofibromatosis type 1, subarachnoid hemorrhage, endovascular treatment

Introduction

Vertebral arteriovenous fistula (VAVF), which can cause subarachnoid hemorrhage (SAH) when having perimedullary drainage,1) is one vascular abnormality reported in patients with neurofibromatosis type 1 (NF-1).1–6) The VAVF with an extracranial vertebral aneurysm (EVAn) in patients with NF-1 has been reported, but is considered to be rare.1–4,6) Furthermore, there is only one reported case of SAH in a case of VAVF with an EVAn, which accompanied with cervical meningocele.1) We report a case of SAH from VAVF with an EVAn in a case of NF-1 without any other lesions or perimedullary drainage.

Case Report

A 59-year-old patient was admitted to our hospital. The patient did not have any previous medical history, but there was a family history of NF-1. The patient was in a comatose state. Diffuse skin lesions were observed on the trunk (Fig. 1A). The patient was intubated and sedated. Computed tomography (CT) revealed SAH (Fig. 1B) and acute hydrocephalus. The CT angiography did not show any apparent intracranial aneurysm, but an extracranial aneurysm with a varix was detected in the left vertebral artery (VA) at the level of C4. The varix was located in the epidural space without perimedullary vein drainage. Subarachnoid hemorrhage was also confirmed at the same level of location of the EVAn and/or varix (Figs. 1C and 1D). The EVAn and varix were considered to be the origin of the SAH. Emergent external ventricular drainage was performed for acute hydrocephalus, followed by cerebral angiography, which showed a left VAVF accompanying the EVAn and varix, and the fistula point was located at the level of C4. A drainage route consisting of the epidural and paravertebral veins were clearly seen, but there was no involvement of any retrograde perimedullary veins such as anterior spinal vein or posterior spinal vein (Figs. 2A–2D). The left VA seemed to be sacrificial as the right VA developed well. Parent artery occlusion

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was performed with endovascular treatment, and following occlusion of the left VA from C3 to C5, the left VAVF with an EVAn and epidural varix resolved (Fig. 2E). There was no apparent supply from the right VA to the left VAVF with an EVAn. After surgical treatment, the patient was referred to a dermatologist for the skin lesions. Histopathological evaluation of the dermal lesions was not performed due to the postoperative state, and we made a clinical diagnosis of NF-1. Tracheostomy was performed 5 days after coil embolization. Rerupture from the left VAVF was not apparently observed with follow-up CT images. The neurological state of the patient did not improve. The patient ended up spending 2 months in our hospital and then was transferred to another hospital. The patient died because of pneumonitis 1 year after the discharge.

Discussion

The VAVF has been reported as a relatively rare vascular abnormality in patients of NF-1.\textsuperscript{1-6} Complication in such cases with an EVAn is considered to be a further rarity.\textsuperscript{1-4,6}

Deans et al. speculated two possible mechanisms about the manifestation of VAVF in patients with NF-1: a congenital lesion or a secondary lesion.\textsuperscript{7} Because of our patient is an adult, we considered the VAVF unlikely to be a congenital lesion, and most likely resulting from a VA with a weakened arterial wall related to NF-1. This hypothesized mechanism can be supported by the coexistence of an EVAn and epidural varix.

Clinical symptoms caused by VAVF with EVAn are variable,\textsuperscript{1-6} but SAH related to VAVF with an
EVAn in a patient of NF-1 observed on radiology has been reported only in one case. In this case, there were limited findings of SAH on MRI, and the final diagnosis was made with a lumbar tap as described by Morvan et al. In the same case, a meningocele was located close to a vertebral aneurysm, and the authors hypothesized that SAH was probably caused by rupture of the EVAn to the meningocele. In our case, no other lesion accompanied the VAVF with a vertebral aneurysm. Our case is considered to be the first case of VAVF with an EVAn causing SAH without any other lesions.

Craniovertebral arteriovenous fistula (AVF) or perimedullary AVF is considered to be a cause of SAH due to perimedullary drainage. The SAH occurred in our case, however, only the vertebral venous plexus was functioning as a drainage route of the VAVF. Even though the involvement of any retrograde perimedullary veins was not confirmed on cerebral angiography, hemodynamic stress to the epidural varix through the VAVF could result in SAH in our case. Intracranial venous congestion related to the VAVF also might cause SAH. In addition to hemodynamic mechanism, tissue fragility of the dura related to NF-1 could cause SAH in our case. The EVAn and epidural varix were located close to the dura mater. Hemorrhage from the epidural varix could have flowed through the dura mater into the subarachnoid space and resulting in SAH. Our speculation on the mechanism of SAH related to rupture of the VAVF with an EVAn and epidural varix seems possible, because the fragility of the dura mater in patients with NF-1 has been described. As SAH was also confirmed at the level of C4–C5 where the EVAn and epidural varix existed, our speculation seems to be applicable.

The treatment of the VAVF with a vertebral aneurysm by direct surgery or endovascular coil embolization has been reported. In a case of Guzel et al., VAVF with a vertebral aneurysm was successfully treated by direct surgery; however, an anterior fixation was needed for postoperative cervical instability. As the pathology of VAVF is a shunt between the arteries and the veins, it can be difficult to control intraoperative bleeding if direct surgery is chosen. Endovascular embolization did not contribute to good outcome in our case probably due to preoperative damage related to SAH. However, to avoid intraoperative bleeding and possible additional surgery, endovascular embolization can be better as the treatment for VAVF with a vertebral aneurysm in the case that the VA of the lesion side can be sacrificed.

Conclusion

We reported a case of NF-1 accompanied with VAVF and EVAn, which was treated with endovascular coil embolization. Possibly due to hemodynamic stress of the VAVF and tissue fragility of the dura related to NF-1, SAH occurred in our case despite the VAVF with an EVAn was lacking of perimedullary drainage.

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Compliance with Ethical Standards and Informed Consent

The authors obtained informed consent from the patient and her family.

Conflicts of Interest Disclosure

The authors report no conflict of interest concerning the findings specified in this paper.

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