Oculomotor Nerve Palsy Associated with Rupture of Middle Cerebral Artery Aneurysm

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Oculomotor nerve palsy (ONP) with subarachnoid hemorrhage (SAH) occurs usually when oculomotor nerve is compressed by growing or budding of posterior communicating artery (PcoA) aneurysm. Midbrain injury, increased intracranial pressure (ICP), or uncal herniation may also cause it. We report herein a rare case of ONP associated with SAH which was caused by middle cerebral artery (MCA) bifurcation aneurysm rupture. A 58-year-old woman with clear consciousness suffered from headache and sudden onset of unilateral ONP. Computed tomography showed SAH caused by the rupture of MCA aneurysm. The unilateral ONP was not associated with midbrain injury, increased ICP, or uncal herniation. The patient was treated with coil embolization, and the signs of oculomotor nerve palsy completely resolved after a few days. We suggest that bloody jet flow from the rupture of distant aneurysm other than PcoA aneurysm may also be considered as a cause of sudden unilateral ONP in patients with SAH.

KEY WORDS: Oculomotor nerve palsy - Middle cerebral artery aneurysm - Subarachnoid hemorrhage.

INTRODUCTION

Unilateral oculomotor nerve palsy (ONP) associated with subarachnoid hemorrhage (SAH) is a clinical manifestation due to direct compression of the internal carotid-posterior communicating artery (IC-PcoA) junction aneurysm6,8. Midbrain injury or uncal herniation associated with increased intracranial pressure (ICP) could also be the causes of ONPs indirectly from the rupture of distant aneurysm other than PcoA aneurysm6.

We report herein a rare case of ONP associated with SAH which was caused by a direct injury by the bloody jet flow from the rupture of middle cerebral artery (MCA) bifurcation aneurysm without usual causes.

CASE REPORT

A 58-year-old woman was admitted to our hospital because of mild headache without any loss of consciousness. There was no history of diabetes mellitus, hypertension, heart disease, stroke, or cancer. On admission, the patient showed no neurological deficits. Computed tomography (CT) revealed only moderate SAH, and left MCA bifurcation aneurysm was found on CT angiography. After a few hours, the patient suddenly complained of prosis and diplopia. The left pupil was 5 mm in diameter and non-reactive to light. Repeated CT revealed increased SAH on the left sylvian fissure, which was suspected of rebleeding. However, there were no radiological evidence of midbrain injury and uncal herniation on the brain CT scan (Fig. 1). In addition, the patient was alert without any neurological deficit except ONP. Immediately, the patient underwent 4-vessel angiography, and only the left MCA bifurcation aneurysm was found with no other aneurysms elsewhere (Fig. 2). The aneurysm was treated with coil embolization immediately after the 4-vessel angiography (Fig. 3). On the first day after the embolization of the aneurysm, the left pupil became isocoric. The proxis was partially improved and the diplopia was disappeared on day 3. Magnetic resonance imaging (MRI) on day 3 showed no abnormal lesion in the midbrain or basal cistern (Fig. 4). Although
the patient had motor dysphasia due to vasospasm on day 6, the ptosis disappeared completely on day 10. The patient was discharged on day 21 and did not have any neurological deficit during 6-month follow-up.

DISCUSSION

ONP is one of the manifestations of intracranial aneurysms and SAH, and IC-PcoA aneurysm is the most common cause of ONP (34-56%). However, it is also possible that the rupture of distant aneurysms or anomalous vessel may cause ONP. Aiba and Fukuda reported anomaly of basilar arteries, posterior cerebral arteries, and superior cerebellar arteries as well as posterior cerebral artery and superior cerebellar artery inversion might cause ONP.

Fox classified the mechanisms of ONP with an intracranial aneurysm: midbrain hemorrhage, direct local pressure by aneurysms, and hemorrhagic dissection of the nerve were considered as a direct cause, whereas increased ICP (from clot, edema, or hydrocephalus) which results in uncinal herniation was classified as an indirect cause.

In the present case, however, it is very unusual because MCA bifurcation aneurysm was not in the vicinity of the oculomotor nerve. It is highly likely the ONP was induced by the bloody jet flow due to rupture of MCA bifurcation aneurysm. Saito et al. reported a case of ruptured ICA aneurysm with contralateral ONP and suggested that bloody jet flow from the rupture of distant aneurysm could rarely induce ONP.

In the present study, we speculate possible a mechanism of ONP in our patient as bloody jet flow from the rupture of distant aneurysm. First, initial and repeated CT revealed no evidence of midbrain hemorrhage, basilar artery anomaly, hemorrhagic dissection, and uncinal herniation with ICP. MRI also revealed no evidence of abnormality of midbrain and basal cisterns, indicating direct cause of ONP being excluded. Second, the patient had short M1 (9 mm from internal carotid artery bifurcation to middle cerebral artery bifurcation), indicating that the aneurysm was closer to the nerve than normal length of M1 (Fig. 2). Third, the direction of the bloody jet flow from the second rupture (bleeding) might have been changed inferiorly to the way of the oculomotor nerve beside PcoA because of the superiorly placed blood clot from the previous rupture. Fourth, clinical symptom of the patient was correlated with the extrinsic cause, indicating typical ONP (unilateral pupil dilation, ptosis, and partial extraocular palsies). Considering the location of the aneurysm, the bloody jet flow could have direction on superomedial portion of the nerve, resulting in typical ONP. Kuromawa et al. described that the anato-
mical relationship between the exit of the PcoA and the entry of the oculomotor nerve into the cavernous sinus that would usually result in supro medial compression of the nerve by the aneurysm. Typical ONP in this situation may manifest as unilateral pupil dilation, ptosis, incomplete or partial extraocular palsies, and oculomotor synkinesis. Finally, the ONP of the present case was recovered completely after 10 days, which is quite unusual and not easy to explain. ONP is usually recovered very slowly, ranging from a few weeks to months.\(^8,11\) Chen et al.\(^9\) reported that the time to complete resolution of ONP takes about 6 months. Furthermore, early treatment might contribute to early recovery. Indeed, Leivo et al.\(^9\) recommended early treatment for aneurysm-induced third nerve palsy, preferably within 3 days, to avoid functionally and cosmetically unwanted disability.

These mechanisms may explain a rare case of ONP associated with SAH which was caused by a direct injury by the bloody jet flow from the rupture of middle cerebral artery (MCA) bifurcation aneurysm.

**CONCLUSION**

ONP induced by a ruptured MCA aneurysm is very rare, and it would better be kept in mind that the ONP with SAH may occur by the rupture of distant intracranial aneurysms.

**References**