Case Report

Hemifacial spasm caused by the brainstem developmental venous anomaly: A case report and review of the literature

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ABSTRACT

Background: Hemifacial spasm (HFS) is usually caused by vascular compression of the root exit zone (REZ) of the facial nerve. Dual compression of the REZ by veins and arteries is also associated with HFS, but venous origin alone is rarely reported. We present a rare case of HFS caused by the brainstem developmental venous anomaly (DVA) treated with microvascular decompression (MVD).

Case Description: A 30-year-old woman presented with the left-sided HFS since the age of 18 years. The brainstem DVA was diagnosed by magnetic resonance imaging (MRI) and followed by two attempts of MVD at other clinics without any improvement. At our hospital, MVD was performed through a left retromastoid craniotomy. Intraoperatively, after detaching the strong adhesions between the cerebellar hemisphere, petrosal dura and lower cranial nerves, and removing the Teflon sponge inserted during the previous operations, the compressing large vein was found, separated from facial nerve REZ and MVD was completed. The postoperative computed tomography angiography and MRI showed the thrombosis of the main trunk of DVA and decompression of the facial nerve REZ. Complete cessation of HFS with hearing preservation was observed with only slight weakness of mimic muscles which disappeared within 3 months after surgery.

Conclusion: HFS associated with brainstem DVA is a very rare condition. MVD of the facial nerve REZ with transposition of the large draining vein should be considered as an effective treatment option.

Keywords: Brainstem, Developmental venous anomaly, Hemifacial spasm, Microvascular decompression, Venous angioma

INTRODUCTION

Hemifacial spasm (HFS) is usually caused by vascular compression of the root exit zone (REZ) of the facial nerve. The offending vessels in most cases of HFS are the anterior and posterior inferior cerebellar arteries and the tortuous vertebrobasilar artery. Dual compression of the REZ by veins and arteries is also associated with HFS, but venous origin alone is rarely reported. There are only few cases described the HFS associated with the brainstem developmental venous anomaly (DVA), also known as cerebral venous angioma. We present a rare case of HFS caused by the brainstem DVA treated with microvascular decompression (MVD).
CASE DESCRIPTION

A 30-year-old otherwise healthy woman presented with left-sided HFS since the age of 18 years. She initially experienced spontaneous intermittent twitching of orbicularis oculi muscle that gradually progressed in frequency and severity with following spread to other muscles innervated by the facial nerve.

A brainstem DVA was diagnosed by magnetic resonance imaging (MRI) revealing the large venous channel near the facial nerve REZ. At the age of 26, a left retromastoid craniotomy was performed with MVD of the facial nerve without any improvement of patient's spasm. One year later, she underwent second MVD at another clinic, but symptoms were unchanged. At admission, the patient presented with severe left clonic-tonic mimic muscles and platysma contractions exacerbated by voluntary facial movements, fatigue, and psychological stress without any acoustic phenomena or other neurological signs.

Preoperative computed tomography (CT) angiography showed the collection of abnormal dilated medullary veins in the left cerebellar peduncle and the brainstem merging into a single large vein. This collecting vein was located in the left cerebellopontine angle and drained into the inferior petrosal sinus. MRI with contrast enhancement demonstrated the typical appearance of the brainstem DVA. The small venous channels forming “caput medusae” were located in the left cerebellar hemisphere adjacent to the fourth ventricle, in the medulla oblongata and the pons. The large draining vein was passing through the brainstem near lateral recess of the fourth ventricle into the cerebellopontine cistern. The vein reached the surface of the brainstem at the root of the facial nerve [Figure 1].

MVD was performed through a left retromastoid craniotomy, with patient in semi-sitting position. Intraoperative observation revealed strong adhesions between the cerebellar hemisphere, petrosal dura, and lower cranial nerves. After detaching of adhesions, the brainstem and the nerve roots were clearly visualized. The Teflon sponge inserted during previous operations was found under the cisternal portion of the facial and the vestibulocochlear nerves and was removed. The large vein emerged the middle cerebellar peduncle and the choroid plexus of the lateral recess of the fourth ventricle dorsal of the origin of the caudal cranial nerves and run through the cerebellopontine cistern under the facial nerve. The facial nerve REZ was cross-compressed with the main trunk of DVA strongly adherent to the pial surface of the brainstem. The small distal branch of DVA was coagulated and cut and the main trunk was occluded near the cerebellar peduncle with temporary clip. The large vein emptied and became flaccid and the vessel was sharply dissected and carefully transposed ventrally to relieve the compression of the facial nerve REZ. The MVD was completed by insertion of Surgicel and fibrin glue between the brainstem and the displaced draining vein, and temporary clip was released [Figure 2]. The postoperative course was uneventful. CT angiography and MRI showed the thrombosis of the main trunk of the brainstem DVA and decompression of the facial nerve REZ [Figure 3]. Complete cessation of HFS with hearing preservation was observed and slight weakness of mimic muscles (House-Brackmann Grade 2) had disappeared within 3 months after surgery.

DISCUSSION

The most common vascular anomalies of the brain are DVA, affecting 2.5–8.7% of population, and represent congenital variant of the normal venous drainage. DVA is composed of dilated medullary veins forming the so-called “caput medusae” and merge in large transcortical or subependymal collector vein. They are considered as compensatory venous pathways developing anomaly, caused by the absence of normal veins because of aplasia, hypoplasia, or occlusion during fetal life. In the posterior cranial fossa, DVA usually drains through the central pontine vein or vein of the lateral recess of the fourth ventricle and empties into petrosal sinus. Most DVAs are asymptomatic and incidentally found during neuroimaging studies. DVA management remains controversial and generally do not require treatment. Although it is generally accepted that DVAs have a very low risk of rupture and bleeding, intracranial hemorrhage may be the present in a small percentage of patients, especially in cases with coincident adjacent cavernous

Figure 1: Preoperative images. (a) Three-dimensional computed tomography angiogram and (b-d) magnetic resonance imaging showing the collection of abnormal dilated medullary veins in the left cerebellar peduncle and the brainstem merging into a single large vein compressing the facial nerve root exit zone.
angioma. Neurological manifestations can be associated with intracerebral clot formation and edema resulting from thrombosis of the DVA. Pathophysiological mechanisms underlying the clinical manifestations of symptomatic DVAs are divided into idiopathic, flow-related (misbalance of the in- and outflow of blood), and mechanical causes (direct compression of neural structures).

Rare cases of DVA causing mechanical compression of the aqueduct with obstructive hydrocephalus were described and treated with ventriculoperitoneal shunting, aqueductal stenting, and endoscopic third ventriculostomy. The large venous collector of a DVA can compress the cranial nerves and cause different clinical presentations. One case of tinnitus was reported and MRI disclosed the compression of the acousticofacial bundle in cerebellopontine angle by dilated draining vein of the brainstem DVA. Direct cross-compression of the trigeminal nerve root entry zone was described in patients with trigeminal neuralgia managed with medication and treated surgically with MVD and percutaneous procedures.

Venous compression of the root of the facial nerve is rare in HFS and the veins alone as offending vessels are reported in 0%–7.9%. It was shown that cure rates in HFS and trigeminal neuralgia patients who had arterial and venous compressions were significantly lower compared to those with arterial compression alone. Jannetta first described small arteriole in contact and a minute venule crossed the caudal aspect of the facial nerve REZ. The arteriole was mobilized and held away with shredded Teflon, and the venule considered as true causative vessel was coagulated and divided, and patient’s HFS slowly disappeared. Wang et al. analyzed 15 cases of HFS caused by venous compression (13 of them underwent repeat MVD) and the facial nerve REZ was decompressed by Teflon or veins were sacrificed. They stressed that women with platysmal involvement and tonic contractions of facial muscles seem to have a vein as an offending vessel.

There are very few cases of HFS associated with the brainstem DVA reported in literature. Chiaramonte et al. presented a man with the left-sided HFS and MRI demonstrated the typical brainstem DVA with “caput medusae” drained through dilated venous trunk into the superior petrosal sinus. The main venous collector went along the edge of the ventrolateral brainstem causing compression of the facial nerve REZ. Surgery was refused and the patient was managed with carbamazepine. Rizek et al. described a woman suffering of the left-sided HFS with tonic contractions of the left face and anterior neck. MRI showed a DVA in the left pons with the large draining vessel in the cerebellopontine angle near the facial nerve REZ. The patient was treated with...
botulinum toxin injections in the affected facial muscles. Arita et al. reported a case of the left-sided HFS caused by the brainstem cavernous angioma with associated DVA. MRI imaging revealed pontine cavernoma and intra-axial major vein existed ventrally to the facial nerve and drained into the inferior petrosal sinus. Surgical exploration of the cerebellopontine angle did not demonstrate vascular compression of the facial nerve REZ. A large vein was found ventral to the facial nerve and the anterior inferior cerebellar artery loop without contacting with the REZ was displaced to separate in from the facial nerve. Postoperatively, the HFS persisted without any change and it was concluded that in this patient, cavernous angioma may compress the facial nucleus and more likely cause of HFS. Only one case of the HFS caused by the brainstem DVA was successfully treated with MVD. In a 53-year-old woman, Chen et al. demonstrated compression of the left facial nerve REZ by distal portion of the parenchymal segment of the brainstem DVA drained into sigmoid sinus. The offending vessel was located between the facial and auditory nerves. The vein was dissected away from the facial nerve REZ and separated with shredded Teflon. The patient’s HFS completely disappeared 1 week after surgery without complications.

Surgical experience gained in the brainstem DVA associated with the trigeminal neuralgia showed good long-term effectiveness of MVD. It was demonstrated that transposition of the DVAs compressing veins and offending arteries is the best optional treatment for trigeminal neuralgia. Ligation of the main trunk of the DVA in four cases was uneventful and patients became pain free, but one patient died due to cerebellar infarction. Any draining vein injury caused by the surgical procedure may result in congestive ischemic damage of the brainstem or cerebellar infarction. Therefore, vascular injuries to the main collector and tributaries of the DVA must be avoided.

In the present case of HFS, postoperative thrombosis of the main venous collector most likely was developed because of temporary clipping and coagulation of the small branch of the brainstem DVA. These surgical manipulations were performed for transposition of the big venous channel away from the facial nerve REZ. Thrombosed draining vein was clearly demonstrated with MRI and CT and had no neurological consequences. Mobilization of the large vein strongly adherent with the pial surface of the brainstem, especially in distal parenchymatous segment of the brainstem DVA, required the use of sharp dissection resulted in damage of the facial nerve root fibers emerging from the pons. Postoperative slight facial nerve palsy was temporary with subsequent full recovery without recurrence of HFS.

CONCLUSION

HFS associated with the brainstem DVA is a very rare condition. MVD of the facial nerve REZ with transposition of the large draining vein should be considered as an effective treatment option.

Declaration of patient consent

Patient’s consent not obtained as patient’s identity is not disclosed or compromised.

Financial support and sponsorship

Nil.

Conflicts of interest

There are no conflicts of interest.

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