Transvenous embolization in spontaneous direct carotid-cavernous fistula in childhood

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ABSTRACT

Carotid cavernous fistula (CCF) is an abnormal arteriovenous communication in the cavernous sinus. Direct CCF results from a tear in the intracavernous carotid artery. Typically, it has a high flow and usually presents with oculo-orbital venous congestive features such as exophthalmos, chemosis, and sometimes oculomotor or abducens cranial nerve palsy. Indirect CCF generally occurs spontaneously with subtle signs. We report a rare case of spontaneous direct CCF in childhood who did not have the usual history of craniofacial trauma or connective tissue disorder but presented with progressive chemosis and exophthalmos of the right eye. This report aims also to describe the safety and success of transvenous embolization with coils of the superior ophthalmic vein and cavernous sinus through the inferior petrosal sinus.

Key words: Childhood, direct carotid cavernous fistula, transvenous embolization

Introduction

The two main types of fistulas in the region of cavernous sinus are direct arteriovenous (AV) fistula, a true carotid cavernous fistula (CCF), and dural AV fistula. The latter is located in the cavernous sinus which has a gradual onset and usually improves spontaneously. Direct CCF is commonly caused by craniofacial injury and in majority of cases the AV communication is a single hole. Spontaneous CCF may also develop with rupture of an intracavernous aneurysm or where there is vessel fragility in association with atherosclerosis, so-called Ehlers-Danlos syndrome¹ or fibromuscular dysplasia, which was ruled out in this patient. There have been few reports of the occurrence of CCF in children. Signs and symptoms and natural history though have been observed to be similar in adults.²-⁴ Congenital CCF has been reported to be treated successfully with endovascular occlusion.⁵ However, two case reports in childhood presented a benign course, which were managed conservatively by ocular control. Doppler examination of the superior ophthalmic vein (SOV) and follow-up angiography showed spontaneous closure of the fistula.²,³ Blood flow through the shunt is usually fast, especially when the hole is large and spontaneous closure is rare. The preferred treatment has been endovascular occlusion procedure. Although the mainstay of treatment of the direct cavernous fistula is transarterial detachable balloon embolization of the affected cavernous sinus, transvenous embolization using coils has recently been applied with increasing acceptability because of its safety and high success rates.

Case Report

A 3-year-old male was brought in the outpatient clinic because of hyperemia and progressive proptosis of the right eye for one month. There was no history of head trauma and no family history of connective tissue disease. He has no symptoms of increased intracranial pressure or signs of neurologic or cranial nerve deficits. An initial noncontrast computed tomography scan showed a dilated right SOV and exophthalmos of the right orbit, a finding that prompted a referral to the neuroendovascular service. Initial workup was a right transfemoral arterial angiography under general anesthesia. The right internal carotid angiography showed a direct type of CCF at the level of C4 and the draining veins are the SOV anteriorly and the inferior petrosal sinus (IPS) posteriorly [Figure 1a and b]. There was no cortical venous reflux. Patient is negative for any connective tissue disorder.
after a genetic analysis. Due to the progressive ophthalmologic problem, the patient underwent coil embolization through the transvenous approach.

Endovascular procedure was started aseptically with patient under general anesthetics. A 4F sheath was inserted through the right transfemoral artery to the right internal carotid artery (ICA) to obtain a baseline angiography. Roadmap images were done during the procedure. Injections into the contralateral ICA and vertebral artery were done to check the retrograde filling of the carotid siphon through the anterior communicating artery and posterior communicating artery and showed a poor collateral flow of the right ICA. Activating clotting time was elevated by intravenously infusing 1000 units heparin. Another 4F catheter was inserted through the left transfemoral vein going to the right internal jugular vein (IJV) into the right IPS, then into the cavernous sinus, and eventually into the right SOV using a Prowler-plus micro catheter (Cordis) with GT 0.012, 45° angle microguide wire. A total length of 276 cm from 28 coils were detached and densely packed from the right SOV to the cavernous sinus. Post-embolization angiography and plain skull X-rays showed dense coiling and complete disappearance of AV shunts, which signifies total occlusion of the fistula and preservation of the patency of the right ICA [Figures 2a and b, 3a and b]. The vital signs were stable in the entire duration of the procedure. He was monitored in the neurosurgery intensive care unit for 24 hours post-embolization. A gradual improvement of the chemosis on the right eye was noted and was discharged after a week without complication. On one-month follow-up, exophthalmos and chemosis on the right eye have completely resolved.

**Discussion**

Indeed, there have been few reports of CCF in children. Although the preferred treatment has been endovascular
occlusion procedure, transvenous embolization using coils has recently been applied with increasing acceptability because of proven safety and high success rates.

It is preferable to wait for 1 to 2 weeks from the clinical onset of the fistula before doing the endovascular closure, because after a recent injury, the carotid artery may have a greater reactivity to endovascular navigation and is more at risk of damage to the wall. However, some fistulas require treatment at the acute phase when they manifest clinical and angiographic features of venous hypertension within the draining leptomeningeal and cortical veins, for these have the higher risk for cerebral hemorrhage. Direct spontaneous CCF in our patient has high flow based on Barrow’s classification. This type of fistula usually presents with ocular-venous congestive features such as exophthalmos and chemosis, and cephalic bruit.

The goal of endovascular treatment is to completely occlude the fistula but preserve the patency of ICA. The agents used for this procedure include detachable coils, liquid embolic agents, and detachable balloons or non-covered or covered stents delivered transarterially or transvenously. The traditional approach has been transarterial embolization with detachable balloons which was first reported in 1978, but this method is currently not being used in Japan. Transvenous embolization, on the other hand, has been accepted as the preferred treatment for CCF. The transvenous route usually involves a retrograde or posterior path through the IJV, and the IPS up to the cavernous sinus. According to one article, the IPS route is the most direct approach and relatively safe, but excessively tight packing of CCF can result in transient oculomotor or abducens nerve palsies and occlusion of the ICA. Protrusion of coils in the ICA with a large fistula can also occur and may cause occlusion as well, but protrusion may be prevented by placement of a non-detachable balloon along the fistula in the ICA. Other complications include subarachnoid hemorrhage if the IPS is injured. Another option, if the IPS is occluded or absent, is an anterior approach through the SOV through the facial vein or a direct surgical access to the SOV, but in the authors’ opinion, this method is not applicable to a pediatric patient. Occlusion of the arterial segment with fistula that is not amenable to endovascular occlusion was also described in one article.

In conclusion, this report has described a rare occurrence of spontaneous direct CCF in childhood managed with transvenous approach through the IPS. Most importantly, the endovascular treatment employed has proven to be safe and effective.

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