De novo formation of a large cavernoma associated with a congenital torcular dural arteriovenous fistula: case report

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The authors report a case of a developmentally normal child with a congenital complex torcular dural arteriovenous fistula (DAVF) who later, in his teenage years, developed several vermian cavernomas within a large cerebellar developmental venous anomaly (DVA). The patient had initially presented with an abnormally large head circumference but no neurological deficits. He underwent several partial embolization procedures in an attempt to decrease the blood supply of the fistula over the course of 8 years. Nine years following initial presentation, he presented with a fourth ventricular hemorrhage, due to development of a new vermian cavernoma adjacent to a previously known vermian DVA and suffered subsequent mild left-sided hemiataxia from which he later recovered. CT angiographic images demonstrated that the vermian DVA drained into the left transverse sinus, which also drained the torcular arteriovenous fistula. A routine follow-up MRI examination 10 years following initial presentation demonstrated interval development of several large cavernomas in the cerebellum, all within the DVA. The patient had no new symptoms at that time and was neurologically intact. This case report highlights the de novo development of multiple cavernous malformations potentially secondary to DAVF-induced venous congestion in a preexisting DVA.


KEY WORDS  dural arteriovenous fistula; pediatrics; cavernous malformation; vascular disorders

VENOUS congestion of developmental venous anomalies (DVAs) associated with cerebral arteriovenous shunts can result in a variety of manifestations, including venous infarction, hemorrhage, or even de novo cavernoma formation.1–8 Herein we describe a case of a young male who presented with a torcular dural arteriovenous fistula (DAVF) that was associated with a DVA that eventually developed into a cluster of de novo cavernous malformations (CMs).

Case Report

This 6-year-old boy initially was brought to an outside institution for evaluation after his mother noted that he had a larger than normal head circumference. A noncontrast CT scan was performed (Fig. 1), revealing a markedly enlarged torcula. At the time of the initial evaluation, the patient had no cognitive impairment, pulsatile tinnitus, heart failure, or localizing neurological symptoms. He underwent a cerebral angiogram, which demonstrated a complex torcular DAVF that was supplied by bilateral posterior meningeal arteries, middle meningeal arteries, occipital arteries, and arteries of the tentorium cerebelli arising off of the meningohypophyseal trunk (Fig. 2). The fistula had antegrade venous drainage; there was no retrograde cortical venous drainage, but there was evidence of mild venous hypertension, with a pseudophebitic pattern seen in late venous phase angiograms. Multiple transarterial embolizations were performed elsewhere over the course of an 8-year period in an attempt to decrease the blood supply to the high-flow fistula.

Eight years following initial presentation, a follow-up
MRI examination demonstrated persistence of the fistula. Incidentally noted was a DVA that drained the vermis and left cerebellum (Fig. 3A). In retrospect, the cerebellar DVA was faintly visualized in the late venous phase of the initial angiogram obtained when the patient was 6 years old. A CT angiogram obtained at a later date again demonstrated the vermian and left cerebellar DVA, which drained into the left transverse sinus, just distal to the torcular (Fig. 3B). At that time there was no CM associated with the DVA. Nine years following initial presentation, the patient presented with headache and was found to have a new hemorrhage in the fourth ventricle (Fig. 3C). An MRI study performed at that time showed a cluster of CMs that had developed in the cerebellar vermis within the DVA (Fig. 3D). The patient developed a mild left hemiataxia as a result of the hemorrhage but eventually made a near-complete recovery. A repeat cerebral angiogram demonstrated no significant change in the appearance of the torcular DAVF with no substantial change in its venous drainage patterns.

Ten years following initial presentation, the patient underwent a routine MRI follow-up, which demonstrated significant enlargement of the vermian CM and left cerebellar tonsil, all within the region of the patient’s known DVA. Interestingly, the patient had reported no new neuro-
logical symptoms, and neurological examination showed no significant abnormalities (Fig. 3E and F). Specifically, his gait (including tandem gait) and ocular movements were completely normal. Repeat cerebral angiography at this time again demonstrated no substantial change in the appearance of the DAVF and no change in its venous drainage pattern.

Discussion

We report on a patient with a DVA sharing a similar venous drainage pathway with a large torcular DAVF that eventually developed into a cluster of CMs, likely secondary to venous congestion due to the marked venous outflow from the DAVF. Both the DVA and the DAVF were likely congenital in nature and coexisted for over a decade prior to the formation of a cluster of CM.

Many prior studies and case reports have reported on de novo formation of CMs in the setting of DVAs. The presumed mechanism of a DVA resulting in the formation of a CM is elevated venous pressure in the territory of the DVA resulting in a cascade of physiological and signaling events leading to the formation of a CM. Prior studies have reported that CMs associated with DVAs often develop in the distal venous radicles of the DVA rather than within the large central DVA itself. This was seen in our patient, who had multiple CMs in the distal radicles of the DVA while the DVA remained patent.

DVA-associated de novo CM formation in the setting of a DAVF or other cerebral arteriovenous shunting lesion has been reported in prior case reports. In some cases the shunt drains directly into the DVA, while in others the DVA is adjacent to, and shares a common venous drainage pathway with, the arteriovenous shunt. Venous congestion within a DVA associated with an arteriovenous shunt can result in a variety of other neurological manifestations, including venous infarct and hemorrhage. One of the keys to managing DVAs associated with arteriovenous shunts is to maintain the patency of the DVA so as not to impair venous drainage of normal brain parenchyma. These cases, including ours, suggest that indeed venous hypertension may be causally related to the development of a CM in the presence of a DVA, rather than suggesting a purely casual association. This observation may shed some light into the mechanisms underlying the formation of CMs.
of a CM in association with a DVA, factors which so far have remained unknown and purely speculative.

One potential limitation of our case report is the fact that we do not have histopathological confirmation that the lesions within the cerebellar vermis were indeed CMs, rather than evolving hemorrhage. However, one would expect that hemorrhage would become smaller and less complex over time whereas in this case the hemorrhagic lesions within the vermis became larger and more complex over time. In addition, acute hemorrhage would typically be associated with clinical symptoms. Lastly, whether the association between the DVA and the torcular AVF was truly the cause of the CM cannot be definitively established.

In conclusion, this case report highlights the de novo development of multiple CMs secondary to DAVF-induced venous congestion in a preexisting DVA.

References
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