Acute Hydrocephalus in an Adult Patient Secondary to Unruptured Arteriovenous Malformation

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Abstract

Acute hydrocephalus as a result of unruptured arteriovenous malformation (AVM) is a rare phenomenon in the adult population. Patients with AVMs typically present with hemorrhage, seizures, or focal neurologic deficits. Hydrocephalus may result from obstructing the native cerebrospinal fluid drainage by the malformation’s architecture. We report a previously healthy 32 year-old male who initially presented to an optometrist with blurry vision, visual obscurations, and papilledema. A large right frontal cerebral AVM with hydrocephalus and papilledema was confirmed by MRI. Unfortunately localized lesions may result in hydrocephalus as a rare presentation in unruptured AVMs.

Key words: Arteriovenous malformation; Cerebrospinal fluid; Hydrocephalus

Introduction

Arteriovenous malformations (AVM) frequently present with hemorrhage, headaches, seizures, and focal neurologic deficits. However, AVMs have rarely been reported to cause hydrocephalus or pseudotumor cerebri-like symptoms.¹⁻¹² The likelihood of causing obstructive hydrocephalus depends on the location of the nidus and draining veins, as well as flow dynamics. Symptomatology may arise from both the AVM itself (i.e., seizures, hemorrhage, headaches, focal deficits) as well as from hydrocephalus or pseudotumor syndromes (i.e. altered mental status, visual changes, gait disturbances, incontinence). The current study identifies a young male who presented in an unusual fashion for AVMs, with vision changes, headaches, and nausea secondary to hydrocephalus caused by CSF obstruction by a draining vein.

Case Presentation

We present an otherwise healthy 32 year-old right-handed male evaluated at an outside hospital for 3–4 months of mild frontal headaches and blurred vision. He was originally seen by an optometrist with complaints of blurred vision and “gray spots” when moving from seated to standing position. Papilledema was noted on physical examination.

He was referred to an ophthalmologist who confirmed this finding and referred the patient to a Neurologist. Subsequent magnetic resonance imaging (MRI) revealed a large right frontal cerebral AVM measuring 5.6cm x 5.4cm x 4.4cm (Figure 1).
Arterial supply was predominantly from the right middle and anterior cerebral arteries, with primary drainage via a large draining vein measuring up to 8 mm and coursing across the right lateral ventricle. This vein abutted the foramen of Monroe before eventually emptying into the vein of Galen. There was marked dilation of the right lateral ventricle and a 6mm midline shift with secondary dilation of the left lateral ventricle. Therefore it appeared as though this large draining vein was causing obstructive hydrocephalus.

He was transferred to a tertiary care center for neurosurgical evaluation. On arrival he was in no acute distress and grossly neurologically intact, with complaints of nausea. He underwent a computed tomography (CT) angiography of the head and neck, confirming a Spetzler-Martin grade IV (size >6cm, non-eloquent brain, deep drainage) right frontal AVM. The next day the patient underwent a diagnostic cerebral angiogram revealing a small aneurysm at the bifurcation of the right A2 segment of the anterior cerebral artery and a large venous aneurysm along the lateral aspect of the nidus (Figures 2). Multi-staged embolization followed by a surgical resection of the nidus was planned for this patient.

**Figure 1:** Serial axial T2 MRI demonstrating large right frontal AVM with hydrocephalus, demonstrating the path of the large draining vein through the right lateral ventricle, compressing the foramen of Monroe, and into the vein of Galen.

**Figure 2:** (A) Arterial and (B) venous phases of a diagnostic cerebral angiogram in the sagittal plane demonstrating a right frontal AVM (fed by the right middle cerebral artery and right anterior cerebral artery), with a large draining vein.
Discussion

Acute symptomatic hydrocephalus secondary to unruptured AVMs in the adult population remains a very rare phenomenon.\(^{13, 14}\) Multiple mechanisms reported for the etiology of the hydrocephalus, including direct mechanical obstruction from draining vessels or the nidus itself, hydrodynamic disequilibrium, overproduction of CSF, and venous congestion.\(^{9, 12, 15, 16}\)

Our patient originally presented with blurry vision and position-dependent visual obscurations in addition to papilledema. Visual changes due to unruptured AVMs are scarcely reported in the literature. Bayri et al. reported one such patient with visual loss as a result of and unruptured AVM, interestingly with a very similar location and drainage to our case report.\(^4\) Their patient was treated with cerebrospinal fluid (CSF) diversion followed by radiosurgery.

Treatment modalities for hydrocephalus secondary to unruptured AVMs may vary and there are several factors to consider. CSF diversion can be achieved by shunting, endoscopic third ventriculostomy (ETV), septostomy, extraventricular drainage (EVD) during surgical treatment, or close observation if minimally symptomatic. Invasive procedures such as EVD placement or ETV carry additional risk of damage to draining vessels. This can result in intraventricular hemorrhage, especially in the case presented where the large draining vein traversed the right ventricle and compressed the foramen of Monroe. AVM location and drainage may determine the best modality for treatment.

Conclusions

Acute hydrocephalus secondary to unruptured AVMs remains a rare phenomenon in the adult population. Unfortunately localized lesions may result in this uncommon presentation, as with this patient.

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