CASE REPORT

Case Report of an Obstructive Hydrocephalus Caused by an Unruptured Mesencephalic Arteriovenous Malformation in a Boy and a Review of Literature

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Abstract:

Objective:
Arteriovenous malformation (AVM) is the most common form of intracranial vascular malformations in adults. Intracranial pediatric AVMs are rare. AVM located in the vicinity of the brain stem in children are even more rare.

Case report:
This study reports a rare case of acute obstructive hydrocephalus following aqueductal stenosis caused by an unruptured grade IV perimesencephalic arteriovenous malformation. An 11-year-old boy admitted to the hospital with progressive headache, nausea and vomiting throughout a month. A Computerized Tomography (CT) showed an obstructive hydrocephaly. A Magnetic Resonance (MR) imaging revealed a mesencephalic AVM compressing the aqueduct. The patient deteriorated in hours and an emergency ventriculoperitoneal shunting was performed. He did well in the early postoperative period. AVM examined with Digital Subtraction Angiography (DSA) in detail for maintaining the definitive treatment by means of endovascular embolization, microsurgery and stereotactic radiosurgery; but the patient was lost to follow up.

Conclusion:
A Pubmed search revealed 34 cases of hydrocephalus caused by an unruptured AVM in the literature, and only four cases were less than 18 years old with unruptured AVM locating in brain stem or posterior fossa. Although focal neurologic deficit, seizure and headache are the most common symptoms, acute neurologic deterioration due to hydrocephalus may be the presenting symptom in these cases. The decrease in intracranial pressure by changing the flow of cerebrospinal fluid (CSF) via an emergency ventriculoperitoneal (VP) shunting or Endoscopic Third Ventriculostomy (ETV) can be a lifesaving procedure that gives a chance for further treatment modalities.

Keywords: Arteriovenous malformation, Aqueduct, Digital subtraction angiography, Hydrocephalus, Mesencephalon, Ventriculoperitoneal shunt.

1. INTRODUCTION

AVM is the most common form of intracranial vascular malformations in adults. Intracranial pediatric AVMs are rare [1, 2]. AVM located in the vicinity of the brain stem in children are even more rare [1, 3]. Hemorrhage is the most common presentation of intracranial AVMs in children [3]. Focal neurological deficit, seizure and headache are the...
most common symptoms for unruptured AVMs in children [4]. An unruptured AVM presenting with an acutely worsening hydrocephalus is a rare event. We found only 34 cases of hydrocephalus caused by an unruptured AVM in the literature, and only eight of them were less than 18 years old and only 12 cases were locating in midbrain or posterior fossa (Table 1).

Table 1. Literature Review of Hydrocephalus due to Unruptured AVM.

| Author                  | Age & Sex | Localization of AVM               | Treatment of hydrocephalus       |
|-------------------------|-----------|-----------------------------------|----------------------------------|
| Geibprasert et al. [5]  | 16, F     | Thalamus                          | VP shunting                      |
|                         | 17, F     | Buzal ganglia & thalamus          | VP shunting                      |
|                         | 35, M     | Temporoparietal                   | VP shunting                      |
|                         | 39, F     | Frontal /callosum                 | None                             |
|                         | 6, F      | Midbrain                          | VP shunting                      |
|                         | 26, M     | Cerebellum                        | VP shunting                      |
|                         | 55, M     | Cerebellum                        | VP shunting                      |
|                         | 2, M      | Cerebellar vermis                 | None                             |
| Park et al. [6]         | 0, M      | Cerebral hemispheres              | None                             |
| Rodriguez and Molet [7] | 83, M     | Posterior fossa                   | ETV                              |
|                         | 64, F     | Posterior fossa                   | ETV                              |
| Montoya et al. [8]      | 0, F      | Torcula                           | Ventriculojugular shunting       |
|                         | 5, F      | Thalamus                          | VP shunting                      |
| Nozaki et al. [9]       | 0, M      | Midbrain                          | VP shunting                      |
| Mindea et al. [10]      | 55,M      | Parietooccipital and Galenic region| VP shunting                      |
| Millar et al. [13]      | ?, M      | NA                                | NA                               |
|                         | ?, M      | NA                                | NA                               |
|                         | ?, M      | NA                                | NA                               |
|                         | ?, M      | NA                                | NA                               |
|                         | ?, F      | NA                                | NA                               |
| Champeaux et al. [15]   | 54, M     | Thalamic insular & capsular        | ETV                              |
| Ono et al. [16]         | 56, M     | Cerebellar vermis                 | ETV                              |
| Tucker et al. [17]      | 63, M     | Pineal region                     | ETV                              |
| Ebini et al. [18]       | 61, M     | Periventricular                   | None                             |
| Bayri et al. [21]       | 37, M     | Frontal                           | VP shunting                      |
| Rezaee et al. [22]      | NK        | NK                                | ETV                              |
| Pribil et al. [23]      | 20, M     | Fronto-parietal                   | None                             |
| DeFoe et al. [24]       | 24, M     | Mesencephalon                     | VP shunting                      |
| Esparza et al. [25]     | 20, F     | Posterior fossa                   | None                             |
| Hoi & Kerber [26]       | 31, F     | Thalamus                          | VP shunting                      |
| Kurita [27]             | NA        | NA                                | NA                               |
| Liu et al. [28]         | 49, M     | Thalamus                          | Endovascular intervention        |
| Lobato et al. [29]      | 42, F     | Mesencephalon                     | VP shunting                      |
| Pereira et al. [30]     | 41, M     | Pineal region                     | ETV                              |
| Present case            | 11, M     | Perimesencephalic                 | VP shunting                      |

ETV: Endoscopic Third Ventriculostomy, NA: Not Available, NK: Not Known, VP: Ventriculoperitoneal

This study illustrates a case of an unruptured pediatric perimesencephalic AVM that presented with acute neurological deterioration due to acute obstructive hydrocephalus in a child.

2. CASE REPORT

An 11-year-old male from abroad admitted to outpatient clinic with progressive headache, nausea and vomiting which had been presented for a month. Neurologic examinations showed bilateral papillary edema and right-sided central facial nerve palsy. He had been investigated for right sided facial palsy at home and was diagnosed with an AVM located in the perimesencephalic region two years ago. Head CT showed enlargement of lateral and third
ventricles with transependymal cerebrospinal fluid oozing (Fig. 1B). A prompt cranial MR revealed serpiginous signal void structures around the midbrain compatible with an AVM and enlargement of the vein of Galen and internal cerebral veins causing mechanical compression of the aqueduct causing an acute obstructive hydrocephalus caused (Fig. 1A). Cerebral DSA was planned for diagnosis and treatment planning, however, due to rapid deterioration in patient’s consciousness DSA was postponed and emergency right VP shunting was done in the same day. His consciousness improved, headache and vomiting disappeared immediately in the early postoperative period. Control CT scan showed good ventricular decompression (Fig. 1B). DSA was performed via right femoral access two weeks after the operation. Bilateral injection of internal carotid arteries showed enlarged posterior cerebral arteries bilaterally, supplying a diffuse AVM by the perforating arteries of the posterior lateral, medial choroidal and posterior communicating arteries. Venous drainage was into enlarged internal cerebral veins and vein of Galen. There was also some reflux to the deep tentorial and adjacent cortical veins due to high flow. The late capillary and venous phase was prolonged because of venous hypertension (Fig. 1C). The AVM was assessed as grade IV according to the Spetzler-Martin AVM grading system. A multimodal treatment protocol including endovascular embolization and stereotactic radiosurgery to portions of residual AVM was suggested. Patient was discharged on his family’s request as neurologically intact except for a mild right-sided facial palsy and was lost to follow-up.

3. DISCUSSION

3.1. Demographic Features

Hydrocephalus caused by an unruptured AVM may be seen in both gender and in every age but rare less than 18 years [5 - 8]. Only 4 cases of pediatric patients -two female & two male- with hydrocephalus and unruptured AVM locating in midbrain or posterior fossa were reported in the literature [5, 8, 9]. We added the fifth pediatric patient with this study.

3.2. Clinical Signs and Symptoms

Hydrocephalus is mostly expected to be happening after the hemorrhage of an AVM into the ventricular system or subarachnoid space [10]. Hydrocephalus is an uncommon neurologic problem in patients with unruptured AVMs [5].
Presentation of a deeply located intracranial unruptured AVM with acute obstructive hydrocephalus is a rare entity in children. Hemorrhage is usually seen with deep seated AVMs due to deep venous drainage and in unruptured AVMs mostly located in cerebral hemispheres in children [4]. In this case however, the AVM was located around the midbrain and not caused hemorrhage. Hemorrhage, seizure, headache and focal neurologic deficits are common neurologic problems at presentation in children with intracranial AVMs [11, 12]. As hemorrhage is the most common symptom in ruptured AVMs, focal neurologic deficit, seizure and headache commonly seen in unruptured AVMs [4, 13]. Incidental AVMs may also be encountered during childhood deterioration [11]. Focal neurologic deficit can be seen in patients with ruptured or unruptured AVMs, but acute neurologic deterioration due to hydrocephalus with unruptured AVM is a rare entity [7, 10]. The presented case had a cranial nerve deficit for nearly two years, but acute neurological deterioration due to hydrocephalus ensued over a short period of hours. According to the patient’s history of the presented case, an intracranial AVM was detected following facial palsy two years ago, and no new sign or symptom added until one month ago. Although the mechanism -like enlargement as a hemodynamic consequence of increased flow, or stimulated proliferation because of shunt- has been a matter of controversy, the increase in size of AVM has been documented [14]. We do not have any knowledge about the size of previously found AVM, but we can suggest that the occurrence of obstructive hydrocephalus and deterioration of the patient is most probably due to enlargement of the size of mesencephalic AVM.

3.3. Cause of Hydrocephalus

Hydrocephalus can occur as a result of the venous outflow and hemodynamic unbalance [5, 8, 10]; or mechanical obstruction of the ventricles by drainage vein or AVM [5, 7], or by compression of the aqueduct [5, 15 - 17] in unruptured AVM cases. Ebuni et al. [18] suggested that hydrocephalus was result of reflux into periventricular and transmedullary veins instead of mechanical obstruction in their study. Overproduction of cerebrospinal fluid may also be a cause of hydrocephalus in an unruptured choroidal AVM case [19]. In the presented case, the hydrocephalus was thought to be due to mechanical obstruction of the aqueduct by AVM.

3.4. Management of Obstructive Hydrocephalus Due to an AVM

Microsurgical resection [11], endovascular embolization [5, 12], stereotactic radiosurgery [4, 5], and/or multimodal therapy [1] consist of current intracranial AVM treatment. Treatment of brainstem AVMs -especially with high (Grade IV and V) Spetzler-Martin grade, is a challenge [9, 20]. Removal of brain stem AVMs with microsurgical resection is difficult and entangled with a high surgical risk. Since they are located deeply and close to vital structures, radiosurgery also has high risk of adverse effect and hemorrhage during latency periods, and rate of obliteration is relatively low. Preoperative embolization is advised before microsurgical excision for grad IV-V unruptured AVMs locating in brainstem [20]. In case of AVM with large size or locating in eloquent area, conservative treatment may be adopted [5].

Obstructive hydrocephalus due to an AVM can be treated by a VP shunting [5, 18, 21] or ETV [6, 15, 22]. ETV is a technique which would not involve a change between supratentorial and infratentorial pressure relationships [7] and it considered as an advantage for being “shunt free” [15]. Malfunctioning or over drainage may be seen following VP shunting [5]. We chose VP shunting in this case because that procedure was the fastest treatment option in the emergency setting in our conditions [5] to treat obstructive hydrocephalus.

Some authors primarily chose to treat AVM in cases with mild symptoms of obstruction that not required emergency [18, 23].

Patient’s family requested discharge although they were fully informed about the disease and treatment options due to financial concerns and patient was lost to follow up.

CONCLUSION

Presentation of a deeply located intracranial unruptured AVM with acute obstructive hydrocephalus is a rare entity in children. Although focal neurologic deficit, seizure and headache are the most common symptoms, acute neurologic deterioration due to hydrocephalus may be the presenting symptom in these cases. The decrease in intracranial pressure by changing the flow of CSF via an emergency VP shunting or ETV can be a lifesaving procedure that gives a chance for further treatment modalities.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

The study was approved by an internal ethics committee at the specialized arthritis centre and was conducted in
accordance with the recommendations of the Declaration of Helsinki.

HUMAN AND ANIMAL RIGHTS

No Animals were used in this research. All human research procedures followed were in accordance with the ethical standards of the committee responsible for human experimentation (institutional and national), and with the Helsinki Declaration of 1975, as revised in 2008.

CONSENT FOR PUBLICATION

A written informed consent was obtained from the patient when he was enrolled.

CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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Declared none.

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