“Drowning Brain in a Pool of CSF” — A Rare Complication of Periencephalic Subdural Panhygroma following Removal of a Posterior Fossa Tumor

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Posterior fossa tumors are commonly encountered in pediatric age group patients. Most of these tumors present with features of hydrocephalus in the child. Conventionally, these cases are managed by suboccipital craniotomy with decompression of the tumor mass to establish the free flow of cerebrospinal fluid (CSF) across fourth ventricle and aqueduct of Sylvius. Following resection of posterior fossa tumor, appearance of subdural hygroma is a rare phenomenon.

Though few cases of subdural hygroma are reported in literature following foramen magnum decompression in Chiari’s malformation, their appearance following posterior fossa tumor resection is alien to medical literature and limited to only two case reports. Here the authors present a patient with periencephalic subdural panhygroma (PSP) following posterior fossa tumor resection who was successfully treated with a ventriculoperitoneal shunt (VPS) to accomplish a symptomatic and radiologic remission.

Introduction

Periencephalic subdural panhygroma (PSP), following posterior fossa tumor resection (PFTR) is an unusual complication. Post-PFTR clinical improvement, followed by gradual onset nonspecific symptoms of raised intracranial pressure (ICP), points out a possibility toward this unusual complication.

Though the exact mechanism of formation of PSP is unknown, the authors speculate the conversion of previously persisting internal hydrocephalus to postoperative external hydrocephalus in a noncompliant brain due to formation of a huge subdural space freely communicating with major cisterns.

Most of the postoperative subdural hygroma reports are confined to foramen magnum decompression in Chiari’s malformations.¹ ² Here the authors describe a case of subdural hygroma involving both supra- and infratentorial space and proposed a nomenclature of “periencephalic subdural panhygroma” (PSP) that is first of its kind in present-day medical literature.

Because of the paucity of this complication, its management is not standardized and purely driven by clinician’s choice.³ Here the authors propose a suitable nomenclature with possible mechanism of formation and a rational treatment for this rare complication.

Case Description

A 1.5-year-old male child weighing 5 kg was admitted to the department with complaints of irritability, vomiting, decrease in the movements of all four limbs, and multiple episodes of seizure for last 4 months. The baby had normal antenatal, natal, and postnatal history with normal-term vaginal delivery.
On examination, the baby was irritable having emaciated and deprived look (Fig. 1A). On physical examination the head circumference was 54 cm with open and bulged anterior fontanelle. Head control was not achieved. Menace reflex was present. Extraocular movements were full with normal light reflex in both the eyes with no apparent facial weakness. There was no nasal regurgitation of food or nasal intonation of crying sound. On examination of motor system, all four limbs were having power of grade 5/5 with intermittent extensor posturing of lower limbs. Bowel and bladder habit was apparently normal.

Magnetic resonance imaging (MRI) of the brain revealed a well-circumscribed T1-hypointense, T2-hyperintense, with peripheral enhancing cystic mass of size 65 × 58 × 51 cc in the midline posterior fossa region compressing the fourth ventricle and brainstem with gross obstructive hydrocephalus and thinned-out cerebral/cerebellar parenchyma (Fig. 1B–D).

The patient was planned for suboccipital craniotomy with a telovelar approach for excision of this posterior fossa mass. A pinkish white, cystic, well-capsulated, moderately vascular mass was discovered. After complete excision of the mass, aqueduct of Sylvius was left well wide open with pulsating CSF slowly filling the roomy postoperative posterior fossa cavity (Fig. 1E). Dura closed in watertight manner to prevent CSF leak.

In postoperative period, the patient showed significant clinical improvement and put on oral diet. On ninth postoperative day, the child became drowsy multiple episodes of vomiting with one episode of seizure with extensor posturing. He was put on intravenous fluids with antiepileptics and sent for a noncontrast computed tomography (NCCT) scan of the brain in emergency basis, which revealed an intercommunicating subdural hygroma spanning both supra- and infratentorial compartment of the brain with a wide and patent aqueduct (Fig. 1F). This unusual phenomenon tempted the authors to propose a new term “periencephalic subdural panhygroma” (PSP). To relieve the raised ICP, a ventriculoperitoneal shunt (VPS) was done in emergency basis. Post-VPS, a significant improvement in sensorium was observed in the child. Even though the clinical status of the child improved, NCCT of the brain still showed presence of PSP with shunt in situ.

![Fig. 1](https://example.com/figure1.jpg)

**Fig. 1** (A) Preoperative picture of the child. (B) MRI of the brain T1 sequence showing hypointense cystic mass in the midline cerebellar region compressing fourth ventricle with gross hydrocephalus. (C) MRI of the brain T1 contrast, featuring a midline cerebellar peripheral contrast-enhancing cystic mass compressing fourth ventricle. (D) MRI of the brain T2 sequence showing cystic midline cerebellar mass isointense with CSF compressing fourth ventricle. (E) Post–tumor resection view of roomy posterior fossa with visible aqueduct of Sylvius with free-flowing CSF. (F) Postoperative NCCT of the brain showing both supra- and infratentorial subdural hygroma with dilated ventricles and brain parenchyma being compressed both intra- and extra-axially. CSF, cerebrospinal fluid; MRI, magnetic resonance imaging; NCCT, noncontrast computed tomography.
After 1 week the child was discharged with oral medication. The histopathology of the tumor came out to be “hemangioblastoma.”

In the first follow-up after 3 months, the child showed significant weight gain with encouraging social smile. Three-month postoperative NCCT of the brain showed almost completely resolved PSP with shunt in situ (Fig. 2C).

Discussion
Subdural hygroma was first reported in 1916 by Payr, which he described as “meningitis serosa traumatica” that was later redescribed as “subdural hygroma” by Dandy in 1932.

Infantile subdural hygroma of surgical interest represents an extremely heterogenous group of pathology with various etiologic factors such as pneumococcal leptomeningeal infections, late complications of traumatic brain injury, and several neurosurgical procedures.

Subdural hygroma as a complication of post–foramen magnum decompression (FMD) for Chiari’s malformation is a known fact and supported by few medical literatures. The exact pathophysiology of hygromas post-FMD where durotony was not performed is unknown, although it is supposed to be contributed by few microscopic breeches during procedure.

Subdural hygroma in infants mostly presents with features of raised ICP, such as irritability, vomiting, altered sensorium, seizure, and focal neurological deficit due to compressive effect on brain parenchyma.

The mechanism of formation of postoperative subdural hygroma is not fully understood. Here, the authors propose the possible mechanism of periencephalic subdural panhygroma in an infant with postoperative posterior fossa tumor resection as follows:

“Long duration hydrocephalus makes the infantile brain noncompliant. The extra space achieved in this noncompliant infantile brain after resection of a huge tumor mass is not compensated by proper enlargement of brain parenchyma; rather it encourages the formation of a potential subdural space around the brain by the ball-valve mechanism of the arachnoid tear where the CSF enters into the subdural space but can’t escape, leading to formation of a PSP in order to maintain the pressure-volume homeostasis within the rigid skull bone. Later, there is a complete communication between intra- and extra-axial compartments with free-flowing CSF having no actual pressure difference between two compartments.”

Several management protocols of subdural hygroma are described in literature, each having specific advantages and disadvantages. Here, contrary to their expectation, the authors observed that CSF coming out just after durotomy was not in pressure. Therefore, the authors preferred a VPS to a subdural-peritoneal shunt for long-term management. This patient with VPS exhibited good clinical and radiologic outcome in follow-up period.

Conclusion
PSP following resection of a posterior fossa tumor can be rightly described as “a drowning brain in a pool of CSF.” VPS is an effective mode of treatment for these cases. Paucity of similar complications in medical literature makes this report an interesting read.
Table 1  Previous cases of subdural hygroma following posterior fossa surgery

<table>
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<tr>
<th>Authors</th>
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<th>Management</th>
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<tr>
<td>Albuquerque et al</td>
<td>1997</td>
<td>JNS</td>
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<td>Arachnoid cyst rupture</td>
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<td>Ranjan et al</td>
<td>1996</td>
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<td>2005</td>
<td>JNS</td>
<td>2</td>
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<td>Stavrinous et al</td>
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<td>BJN</td>
<td>1</td>
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<td>2009</td>
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<td>Suzuki et al</td>
<td>2011</td>
<td>Neurol Med Chir (Japan)</td>
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<td>Oomman et al</td>
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Abbreviation: VPS, ventriculoperitoneal shunt.

References