CSF hydrothorax without intrathoracic catheter migration in children with ventriculoperitoneal shunt

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Abstract

Background: Thoracic complications of ventriculoperitoneal (VP) shunts have been extensively reported in the literature. Cerebrospinal fluid (CSF) hydrothorax without catheter migration, however, has been rarely described and poorly understood.

Case Description: We describe development of pleural effusion and respiratory distress in a 3-year-old boy with no evidence of VP shunt catheter displacement on plain radiograph and stable ventricle size on rapid sequence magnetic resonance imaging (MRI) brain. Chest X-ray revealed complete opacity of right hemithorax. Pleural effusion was consistent with transudate. Beta-2 transferrin returned positive. The patient underwent externalization of VP shunt, and upon resolution of effusion, re-internalization with new distal shunt catheter. A literature review of CSF hydrothorax in children without intrathoracic shunt migration was performed. Eleven cases were identified in the English literature. Age at VP shunt placement ranged from birth to 8 years of age. Interval from VP shunt placement to CSF hydrothorax ranged from 1.5 months to 5 years. History of shunt revision was reported in two cases. Presenting symptoms also included ascites and inguinal hernia or hydrocele. Reported diagnostic studies consist of CSF culture, radionuclide shuntogram, beta-2 transferrin, and beta-trace protein. Laterality of the VP shunt and development of pleural effusion were predominantly right sided. Definitive surgical treatment included VA shunt, repositioning of the peritoneal catheter, and endoscopic choroid plexus coagulation.

Conclusion: CSF hydrothorax is a rare thoracic complication of VP shunt placement with no radiographic evidence of shunt migration or malfunction. Postulated mechanisms include limited peritoneal capacity to resorb CSF in children and microscopic communications present in congenital diaphragmatic hiatuses.

Key Words: Cerebrospinal fluid hydrothorax, shunt malfunction, ventriculoperitoneal shunt

INTRODUCTION

Complications of ventriculoperitoneal (VP) shunt have been reported extensively in the literature. Thoracic manifestations include pleural effusion, bronchial perforation, pneumothorax, and pneumonia. Cerebrospinal fluid (CSF) pleural effusion in the absence of migration of distal VP shunt catheter in
children, however, has been rarely described and poorly understood[11]. We report a case of CSF hydrothorax in a child with no radiographic evidence of VP shunt migration and review the literature on associated clinical findings.

**CASE HISTORY**

**History and Examination**

The patient is a 3-year-old boy born at 40 weeks by Cesarean section who initially presented with congenital hydrocephalus. He had right VP shunt placement at birth, with subsequent revisions at 13 and 23 months of age. He became symptomatic 10 days prior to presentation with progressive viral-like upper respiratory symptoms, including poor oral intake, intermittent fever, cough, irritability, and an ill appearance. Patient’s co-morbidities include intractable epilepsy and craniosynostosis.

On examination, he was somnolent, but he would open his eyes spontaneously. He had full strength in all extremities. His shunt incisions and shunt track were nonerythematous and nontender. No swelling was seen along the track. He demonstrated intermittent cough with mild desaturations to 89%. Laboratory studies were within normal limits aside from mild thrombocytopenia (79 × 10^3/mcL) and elevated valproic acid level (183 mg/L; reference therapeutic range 50–100 mg/L).

**Imaging**

Chest radiograph revealed complete opacification of the right hemithorax with mediastinal displacement [Figure 1]. Radiographic shunt series revealed radiographically intact VP shunt without migration into the thorax [Figure 2]. Quick-brain magnetic resonance imaging (MRI) revealed stable appearance of the ventricles without change in ventricular size or new extra-axial fluid collections. Ultrasound of the abdomen and the thorax confirmed right sided pleural effusion, underlying atelectatic lung, and a small amount of ascites.

**Hospital course**

A chest tube was inserted on the right side with drainage of 800 ml of straw colored fluid under pressure. The fluid profile was that of a transudate without infection. The patient’s mental status immediately improved. Chest tube output was brisk with 500 ml over the initial 24 h, and increasing to an output of approximately 50 ml/h. The valproic acid dose was decreased given the supratherapeutic level.

Beta-2 transferrin was sent from the chest tube drainage, resulting in a positive study. The final CSF culture was negative for infection. A decision was made to externalize the VP shunt at the abdomen, following which the chest tube output decreased considerably. The chest tube was placed to water seal with no re-accumulation of pleural fluid. The patient returned to the operating room for replacement and internalization of distal peritoneal tubing.

Postoperatively, the chest tube was maintained on water seal for 48 h without an increase in output. Serial ultrasound exam demonstrated no re-accumulation of pleural fluid, and the chest tube was subsequently removed. At 1 year follow-up, he has no re-accumulation of pleural fluid, and no signs or symptoms of shunt malfunction.

**DISCUSSION**

Upon initial presentation, the right chest opacity on chest radiograph was initially thought to be related to valproate toxicity resulting in an eosinophilic effusion. Consistent with this diagnosis was the patient’s depressed mental state and thrombocytopenia. When the chest tube was
inserted, however, pleural fluid analysis was negative for eosinophils. Pleural fluid was transudative with mostly macrophage predominant cellularity and no evidence of malignant cells. The patient did not have symptoms of common etiologies for transudative effusions, such as congestive heart failure (no cardiomegaly on chest X-ray, no S3 on auscultation, no hepatomegaly, normal pulses and blood pressure), nephrotic syndrome (normal urine output, normal serum albumin, no peripheral edema), and liver failure (normal hepatic function tests, normal coagulation tests). Despite improved respiratory status following pleural drainage, the patient continued to have persistent clear fluid drainage from his chest tube at approximately 50 ml/h. Beta-2 transferrin was sent, and its positive result confirmed the diagnosis of CSF hydrothorax.

Thoracic complications of VP shunts have been previously outlined into three categories: intrathoracic trauma during placement of a shunt, migration of the peritoneal catheter into the chest, or pleural effusion accompanying CSF ascites. In the absence of iatrogenic injury or migration of the peritoneal catheter, symptomatic CSF hydrothorax may infrequently result with and without concomitant CSF ascites. Absence of radiographic signs of shunt malfunction, disconnect, or intrathoracic migration of the catheter, however, raises a diagnostic challenge. Radionuclide shuntogram and beta-2 transferrin assays of pleural fluid are of significant diagnostic utility in suspected cases of CSF hydrothorax.

The mechanism of CSF hydrothorax in children without VP shunt catheter displacement remains less clear. Migration of CSF from the peritoneal to the pleural cavity depends on two factors: Malabsorption of CSF in the peritoneal cavity and open communication between the peritoneal and the pleural cavities to enable intraperitoneal CSF to pass into the pleural cavity. A theoretical risk factor is the limited peritoneal capacity to resorb CSF in children, resulting in CSF ascites and associated pleural effusion. Possible contributory factors also include history of abdominal infection, abdominal surgery, and formation of pseudocysts. Another possibility is mechanical leakage of CSF from the shunt valve, the catheter, or between their connection, which is most likely suspected to be the case in our patient.

Conduits for intrathoracic catheter migration traditionally have been suggested to involve congenital diaphragmatic hiatuses, such as the anterior foramen of Morgagni and the posterior foramen of Bochdalek. In children, these areas in the diaphragm are also easily eroded or can harbor microscopic communications undetectable by thoracoscopy or nuclear imaging studies. Chronic inflammation can further facilitate transudation of CSF fluid via capillary and lymphatic channels in the diaphragm. A cyclic pressure gradient, created by negative intrathoracic pressure during inspiration and positive abdominal pressure during expiration, is presumed to allow unidirectional flow of CSF.

In view of these mechanistic factors, a literature review of CSF hydrothorax in children without intrathoracic catheter migration was performed. A total of 11 pediatric cases of CSF hydrothorax without intrathoracic catheter migration were identified in the English literature [Table 1]. While most VP shunts and resultant hydrothorax were left-sided, laterality of VP shunt did not overlap with poor abdominal re-absorptive capacity are susceptible to development of hydrocele or inguinal hernia following placement of VP shunt. While relatively little is known whether these presenting abdominal symptoms have any relation to CSF hydrothorax development, ascites appeared to occur mutually independent from inguinal hernia or hydrocele [Table 1]. Consistent with this trend, findings in our patient included mild ascites but no inguinal hernia or hydrocele.

As reported in Table 1, diagnostic studies reported in the literature, in descending order of frequency, consisted of CSF culture, radionuclide shuntogram, beta-2 transferrin, and beta-trace protein. CSF culture was documented in eight cases (73%). Radionuclide shunt study was obtained in five cases (45%). Beta-2 transferrin was sent in four cases (36%), all of which returned positive. One report utilized beta-trace protein, which was positive. Variability in diagnostic methods may be related to diagnostic preference or institutional availability of these tests as reflected by the wide geographic origin of the report series.

A chest tube was placed in 5 of 11 cases (45%) for management of pleural effusion. Laterality of VP shunt was right-sided (82%), left-sided (9%), and not reported (9%). Pleural effusion developed in the right lung (82%), left lung (9%), and bilateral lungs (9%). While most VP shunts and resultant hydrothorax were right-sided, laterality of VP shunt did not overlap with concomitant hydrothorax when it occurred in the left. There was one case of right-sided VP shunt resulting in bilateral involvement of the lungs.
Table 1: Series of CSF hydrothorax in children without intrathoracic catheter migration

| Series | Origin | Prior shunt revision | Prior abdominal surgery | Ascites | Inguinal hernia or hydrocele | CSF culture | Beta-2 transferrin | Radionuclide shuntogram | Chest tube | Laterality of VP shunt | Laterality of pleural effusion |
|--------|--------|----------------------|-------------------------|---------|-----------------------------|-------------|-------------------|-------------------------|------------|----------------------|-------------------------------|
| Globl and Kaufman (1978)[5] | Philadelphia, PA, USA | No | No | Yes | No | Sterile | NR | No | No | Right | Right |
| Faillace and Garrison (1998)[6] | Jacksonville, FL, USA | Yes | No | No | No | Sterile | NR | Yes | No | Right | Bilateral |
| Hadzikaric et al. (2002)[7] | Al-Koban, Saudi Arabia | No | No | Yes | Sterile | NR | No | Yes | Right | Left |
| Adeolu et al. (2006)[8] | Osun state, Nigeria | No | No | No | NR | NR | No | No | Right | Right |
| Born et al. (2008)[9] | Bonn, Germany | No | Yes | No | No | Sterile | NR* | No | No | Left | Right |
| Smith and Cohen (2009)[10] | Toronto, Canada | No | No | Yes | No | Sterile | Positive | No | Yes | Right | Right |
| Chuen-im et al. (2011)[11] | St. Louis, MO, USA | Yes | Yes | No | Sterile | Positive | Yes | Yes | Right | Right |
| Kocaogullar et al. (2011)[12] | Konya, Turkey | No | No | No | Yes | NR | NR | Yes | Yes | Right | Right |
| Patel et al. (2011)[13] | Little Rock, AR, USA | No | No | Yes | Sterile | Positive | Yes | Yes | Right | Right |
| Ulus et al. (2012)[14] | Samsun, Turkey | No | No | No | NR | Positive | Yes | Yes | Right | Right |
| O’Halloran et al. (2013)[15] | Dublin, Ireland | No | Yes | No | Sterile | NR | No | No | Right | Right |

*Beta-trace protein. *Abbreviation NR (Not Reported)

Final surgical treatment modalities ranged from ventriculo-atrial (VA) shunt placement (73%), reposition of peritoneal catheter (18%), and endoscopic choroid plexus coagulation (9%). Of eight patients in whom VA shunt was eventually placed, the catheter was externalized prior to VA shunt placement in two cases, and the peritoneal catheter was repositioned prior to VA shunt placement in one case. Externalization of a distal catheter may guide subsequent strategy for treatment, whether it is conversion to a VA shunt or replacement of the distal VP shunt catheter. Ventricle-pleural shunt was attempted in one patient but subsequently had to be revised to a VA shunt. Further follow-up data may be useful for determination of long-term efficacy of various surgical revision modalities.

CONCLUSION

CSF hydrothorax is a rare but important complication of VP shunt placement in children without evidence of shunt migration or malformation. Postulated mechanisms include limited peritoneal capacity to resorb CSF in children and microscopic communications present in congenital diaphragmatic hiatuses. In suspected cases, beta-2 transferrin assay and radionuclide tracer shunt series are useful diagnostic studies. CSF culture is commonly obtained to exclude infection. Thoracocentesis of pleural fluid or chest tube placement facilitates management of persistent pleural effusion. Externalization of the distal shunt catheter may resolve the hydrothorax and may guide subsequent treatment strategy.

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