Ruptured pediatric cerebellopontine angle epidermoid cyst: a case report detailing radiographic evolution and clinical course

*Zhe Guan, MS,† Todd Hollon, MD,‡ J. Nicole Bentley, MD,§ and Hugh J. L. Garton, MD, MHS‡

†University of Michigan Medical School; and ‡Department of Neurosurgery, University of Michigan Health System, Ann Arbor, Michigan

Epidermoid cysts (ECs) are uncommon pediatric tumors that often occur in the cerebellopontine angle. Although cyst rupture is a recognized complication, the radiographic evolution of an EC following rupture and the resultant parenchymal brainstem edema have not been reported. The authors present the case of a 13-year-old female with a newly diagnosed cerebellopontine angle EC who presented with worsening headaches, photophobia, and emesis. Magnetic resonance imaging demonstrated significant pericystic brainstem edema and mass effect with effacement of the fourth ventricle. Refractory symptoms prompted repeat imaging, revealing cyst enlargement and dense rim enhancement. Resection of the EC resolved both her symptoms and the brainstem edema. This case documents the radiographic evolution of EC rupture and subsequent clinical course.

http://thejns.org/doi/abs/10.3171/2015.4.PEDS153

KEY WORDS cerebellopontine angle tumor; chemical meningitis; cyst rupture; epidermoid cyst; oncology

Epidermoid cysts (ECs) represent about 1% of all intracranial tumors.\(^{19,28}\) Their incidence is highest in the 3rd to 5th decades of life, and they infrequently occur in pediatric patients. Intracranial ECs are benign congenital tumors that arise from displaced ectoderm during neural tube closure early in embryonic development.\(^{6,19,28}\) About 40%–50% of intracranial ECs are found in the cerebellopontine angle (CPA), with other common sites including the parasellar region, along the tentorium, and the fourth ventricle.\(^{6,10,23}\) Epidermoid cysts constitute 4% of all tumors found in the CPA, making it the third most common CPA tumor behind vestibular schwannomas and meningiomas in children and adults.\(^{11}\) In contrast, dermoid cysts, although similar in pathophysiology with the inclusion of ectopic ectoderm during neural tube closure, are otherwise easily differentiated from ECs by their midline location, radiographic findings consistent with fat rather than CSF, and microscopic features. Additionally, dermoid cysts compose only 0.5% of all intracranial tumors, making them less common.

The natural history of ECs is that of indolent linear growth beginning at birth, with symptoms developing later in life because of neurovascular or brainstem compression. Chronic symptoms such as headache, cranial neuropathies, and cerebellar dysf

Case Report

Clinical Presentation

A 13-year-old right-handed female with a history of...
headaches presented to the pediatric emergency department (ED) with a 3-day severe, atypical occipital headache associated with dizziness. At this time she denied photophobia, emesis, or neck stiffness and had a normal neurological exam. She was initially treated with ketorolac, prochlorperazine, and diphenhydramine and was discharged from the ED with a planned outpatient follow-up with her pediatrician.

Three days later, she returned to the ED with continued severe atypical pain, reporting that the medications provided only transient relief. An MRI study was performed, revealing a 2.8 × 1.3 × 2.5–cm nonenhancing lesion in the right CPA exerting mild mass effect on the adjacent pons. The lesion was characterized by minimal pericystic FLAIR signal (Fig. 1A) or rim enhancement (Fig. 1C), with densely restricted diffusion (Fig. 1B). Given these imaging findings, the patient was diagnosed with an EC. Because she had no focal neurological signs referable to the mass lesion and because MRI was not believed to show an acute process, a migraine headache syndrome was considered to be a likely etiology of her symptoms. The patient was conservatively managed with analgesics and discharged with a recommendation for outpatient follow-up.

Approximately 2 weeks later, she again presented to the ED with refractory symptoms now associated with nausea, emesis, and photophobia, and methylprednisolone was prescribed. Treatment initially relieved her symptoms, but she returned 1 week later with recurrent headaches and nausea, and dihydroergotamine was administered intravenously. Because of the limited response to medi-

**FIG. 1.** Brain FLAIR (upper), diffusion-weighted (center), and T1-weighted (lower) post-Gd sequences demonstrating the progression of brainstem edema and cyst growth following EC rupture. A–C: Initial imaging shows the classic appearance of a CPA EC with mild compression of adjacent brainstem and dense diffusion restriction. D–F: Following cyst rupture, parenchymal edema with mass effect is seen involving the pons and cerebellum leading to distortion of the fourth ventricle. G–I: Resolution of edema with the administration of dexamethasone is seen on FLAIR. Note that the image in panel I is a preoperative MRI study for stereotactic navigation completed approximately 3 weeks after the images in panels G and H.
cal therapy, her diagnosis was reconsidered and a lumbar puncture was performed. This revealed an opening pressure of 31 mm Hg, a white blood cell count of 240/μl (50% lymphocytes, 41% neutrophils), a red blood cell count of 3/μl, protein level of 94 mg/dl, and glucose level of 55 mg/dl. Repeat MRI demonstrated extensive new FLAIR signal abnormalities within the pons and middle cerebellar peduncle, with extension into the right cerebellar hemisphere (Fig. 1D). The lesion was not significantly changed in size and continued to exert mass effect, with distortion of the fourth ventricle. A new thickened enhancing rim suggesting reactive changes was present (Fig. 1F). In view of these clinical findings, EC rupture and chemical meningitis were diagnosed. Since the patient was neurologically intact, she was given a trial of dexamethasone that led to the rapid resolution of her symptoms. She was discharged on hospital Day 5 with a planned outpatient follow-up.

Two weeks later, she returned for an outpatient follow-up, reporting good control of her symptoms. At that time, repeat MRI showed radiographic resolution of the brainstem edema (Fig. 1G). A long steroid taper was initiated but resulted in recurrent headaches and dizziness with accompanying eye discomfort and emesis. Because the EC had not spontaneously resolved and given the intolerable side effects of the steroids, the decision was made to proceed with operative intervention. A thin-slice MRI study was performed for intraoperative navigation and showed significant interval enlargement (2.7 × 2.7 × 3.4 cm) with an avidly contrast-enhancing rim (Fig. 1H and I). Near-complete effacement of the fourth ventricle was noted in response to increased mass effect from the expanding EC following rupture.

Operative Intervention and Pathological Findings

A right retrosigmoid craniotomy was performed, and a pearly white lesion was found deep to and spanning the cranial nerve (CN) complexes of VII–VIII and IX–XI (Fig. 2). From the viewable operative corridor, the capsule of the lesion was not obviously ruptured, and no cyst contents were visibly free in the subarachnoid space. However, only a small portion of the entire capsule was visible. The capsule was opened and caseous material was extracted. Intracapsular gross-total resection was performed, with removal of the tumor in a piecemeal fashion. On inspection, the capsule was quite variable in thickness and strength. As the caseous material was removed, we observed several perforations associated with thin areas of the capsule, although it is possible these were the result of the dissection process. Other portions of the capsule were thicker and densely adherent to the surrounding CNs. These areas were left behind to avoid CN injury. Histological examination of intraoperative tumor specimens showed a cyst capsule composed of stratified squamous epithelium. Cystic contents demonstrated laminated keratin debris, as well as compact keratin components (Fig. 3).

Postoperative Course

The patient tolerated the procedure well and was discharged on postoperative Day 3 without any neurological deficits. At the 3-month follow-up, she remained clinically well and had returned to school without any functional impairment. She reported only mild headaches that were easily treated with ibuprofen. Follow-up MRI at 6 months revealed resolution of the brainstem edema (Fig. 4A) and a small residual capsule. No rim enhancement was noted, and normal fourth ventricular anatomy had been restored (Fig. 4B and C).

Discussion

This case is the first description of the clinical course of a ruptured pediatric CPA EC with corresponding radiographic evolution. Although the capsule did not appear to be open on initial operative inspection, we could only visualize a small portion of the total surface of the lesion.
is thought to be rare, resulting in a clinical presentation consistent with chemical meningitis. Patients with persistent or disabling symptoms related to neurovascular compression or recurrent cyst rupture should be offered surgical intervention. In patients with normal neurological function and no signs of raised intracranial pressure, conservative management with close observation may be recommended initially. Because ECs grow slowly, this approach can make surgery unnecessary or delay surgery by several years. As these lesions are poorly responsive to chemotherapy and radiation, excision remains the definitive treatment. Ideally, gross-total resection is performed to minimize the risk of tumor recurrence. The overall recurrence rate is reported to range from 0% to 36%. The rate of reoperation due to recurrence ranges from 0% to 21%. The natural history of recurrence in these lesions is difficult to ascertain from the literature, as the relatively short follow-up and slow growth pattern limit proper assessment. The prospect of recurrence due to postoperative residual cyst presents a difficult challenge for intraoperative decision making, as the tumor capsule is often densely adherent to neighboring neurovascular structures. Controversy exists between those who advocate for maximum resection at the cost of possible neurological deficit to prevent recurrence and those who favor a more conservative approach, accepting subtotal resection to avoid both transient and permanent neurological injury. Recent data suggest that the total removal of CPA ECs does not significantly increase morbidity or mortality and should be the operative goal, although another series reported similar rates of recurrence for gross-total and subtotal resection (23% and 27%, respectively). Clinical judgment must ultimately be used to determine the most appropriate extent of resection for each patient based on clinical presentation and intraoperative findings.

Regardless of the extent of tumor removal, complications are relatively common. Aseptic meningitis is reported to occur in 10%–40% of cases. This is attributed to intraoperative spillage of cyst contents and can usually be treated successfully with a short course of oral steroids. Transient CN deficits are common following surgery, especially affecting nearby CNs V–VIII. Deficits of CNs V–VII and X tend to be transient and typically resolve over 0.5–1.5 years, but damage to CN VIII with hearing loss often persists. Less common complications include CSF leakage (4%–8%), symptomatic hydrocephalus (14%), vestibular or cerebellar dysfunction, and pseudomeningocele (5%).

Conclusions

Following CPA EC rupture causing pericystic brainstem edema, our patient experienced resolution of her presenting symptoms following subtotal resection without postoperative neurological deficits or complications. This case warrants attention because it is the first report to demonstrate that cyst rupture leads to radiographic findings that challenge the traditional description of a mass lesion without significant parenchymal edema. The documentation of a ruptured EC in evolution provides evidence for the possibility of a more insidious clinical course resulting from these “benign” lesions.
References

Disclosure
The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions
Conception and design: all authors. Acquisition of data: Guan, Hollon, Bentley. Analysis and interpretation of data: all authors. Drafting the article: Guan, Hollon, Bentley. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Garton. Administrative/technical/material support: Garton. Study supervision: Garton.

Correspondence
Hugh J. L. Garton, Department of Neurosurgery, University of Michigan, Taubman Health Care Center, Rm. 3552, 1500 E. Medical Center Dr., Ann Arbor, MI 48109-5338. email: hgarton@med.umich.edu.