A rachnoid cysts are developmental collections of cerebrospinal fluid (CSF) contained within the arachnoid membrane and subarachnoid space. Their anatomical distribution follows that of the main cerebral cisterns.6 These cysts can expand, distort, and compress the surrounding neural structures, eventually producing symptoms.13

Aubry and Ombrédanne, in their first report on cerebellopontine angle (CPA) arachnoid cysts, classified these lesions into 3 groups: pseudotumoral, cystic, and adhesive.2 With progress in the understanding of arachnoid cysts this classification, which is mostly based on lesion size, has fallen out of use.

In most of these cases, the clinical presentation included cerebellar signs and/or endocranial hypertension–related symptoms (e.g., headache, nausea, vomiting) (Table 1). Three pediatric patients with CPA arachnoid cysts, 2 with hearing loss and 1 with recurrent headache, who underwent neurosurgical treatment in our institute prompted us to write this report.

OBJECTIVE Few cases of cerebellopontine angle (CPA) arachnoid cysts in pediatric patients have been described in the literature, and in only 2 of these cases were the patients described as suffering from hearing deficit. In this article, the authors report on 3 pediatric patients with CPA arachnoid cysts (2 with hearing loss and 1 with recurrent headaches) who underwent neurosurgical treatment at the authors’ institution.

METHODS Four pediatric patients were diagnosed with CPA arachnoid cysts at the International Neuroscience Institute during the period from October 2004 through August 2012, and 3 of these patients underwent surgical treatment. The authors describe the patients’ clinical symptoms, the surgical approach, and the results on long-term follow-up.

RESULTS One patient (age 14 years) who presented with headache (without hearing deficit) became asymptomatic after surgical treatment. The other 2 patients who underwent surgical treatment both had hearing loss. One of these children (age 9 years) had recent-onset hypacusia and experienced complete recovery immediately after the surgery. The other (age 6 years) had a longer history (2 years) of progressive hearing loss and showed an interruption of the deficit progression and only mild improvement at the follow-up visit.

CONCLUSIONS CPA arachnoid cysts are uncommon in pediatric patients. The indication and timing of the surgical treatment are fundamental, especially when a hearing deficit is present.

https://thejns.org/doi/abs/10.3171/2017.8.PEDS17341

KEY WORDS cerebellopontine angle; arachnoid cyst; congenital
Methods

During the period from October 2004 through August 2012, CPA arachnoid cysts were diagnosed in 4 pediatric patients at the International Neurosciences Institute in Hannover, and in 3 of these cases, the cysts were treated surgically. The surgically treated patients were 2 girls and 1 boy, aged, respectively, 9, 14, and 6 years. The arachnoid cysts were located in the right CPA in all cases. The fourth case involved an 11-year-old boy who underwent removal of a left temporal low-grade glioma and was found to harbor an asymptomatic right CPA arachnoid cyst. This patient has been followed for 10 years without any change in the size of the cyst, which has remained clinically silent.

Signs and Symptoms

The first surgically treated patient was a 14-year-old boy who suffered from recurrent headaches for 5 months before the diagnosis. The other 2 surgically treated patients had cerebellar signs with progressive hearing loss. In one of these patients, the 9-year-old girl, the CPA arachnoid cyst was detected only 2 weeks after the onset of the deficit, with consequent direct treatment. In contrast, the other patient with hearing loss, a 6-year-old boy, was only referred to our institute 2 years after the beginning of the hearing deficit due to delayed diagnosis.

Radiological Findings

All patients underwent MRI with T1- and T2-weighted, fluid attenuated inversion recovery (FLAIR), and diffusion-weighted sequences. All the cysts appeared as well-defined, nonenhancing lesions isointense to CSF. High-resolution sequences, such as constructive interference in steady state (CISS), helped the precise delineation of cyst wall and adjacent neural and vascular structures. There was no preoperative diagnosis of hydrocephalus. The mean sizes of the lesions in the craniocaudal, lateromedial, and anteroposterior dimensions were 11.8 mm, 16.8 mm, and 13.6 mm, respectively.

Surgery

In all 3 cases, surgery was performed via a retrosigmoid craniotomy with the patient in a semisitting position, after a screening to exclude patent foramen ovale. The dura was incised in a C-shaped fashion. To minimize cerebellar retraction, the lateral cerebellomedullary cistern
Management of CPA arachnoid cysts in children

J Neurosurg Pediatr Volume 21 • February 2018

121

was opened for CSF drainage. After gentle retraction, the posterior wall of the arachnoid cyst was visualized (Fig. 1) and opened. After aspiration of the fluid, the cyst wall was carefully dissected from the surrounding vascular and neural structures in order to obtain extensive removal (Fig. 2). The area caudal and cranial to the seventh and eighth cranial nerve (CN VII-VIII) complex was carefully explored and further intracystic septations were opened and removed, providing a wide communication with the subarachnoid cisterns.

Jugular venous compression was always performed before dura closure to detect any damaged vein that could cause postoperative bleeding. Opened mastoid air cells were carefully occluded with fragments of fat harvested subcutaneously and sealed with fibrin glue. During the entire procedure brainstem auditory evoked potentials were measured to monitor acoustic nerve function and electromyography recordings of the orbicularis oris and oculi muscles were used to monitor facial nerve function.

Results

There were no deaths in this surgical series. The patient with headaches became asymptomatic and remained so throughout the subsequent 12 years of follow-up. The patient with the recent onset of hypacusia experienced complete recovery immediately after the surgery and has remained asymptomatic throughout 13 years of follow-up. The second patient with hearing deficit, who had a longer history (2 years) of progressive hearing loss before the surgical treatment, showed an interruption of the deficit progression and only mild improvement over 5 years of follow-up.

Despite the different postoperative evolution of the symptoms, the follow-up imaging studies performed 6 months after surgery showed a reexpansion of the previously compressed cerebellum (Fig. 3) in all 3 patients with surgically treated CPA arachnoid cysts. At the 1-year follow-up evaluation, MRI demonstrated symmetry of the CPA cisterns bilaterally in all 3 patients with no signs of recurrence.

Discussion

There are few reports in the English literature regarding surgical treatment of CPA arachnoid cysts in children. Including the 3 patients described here, 18 cases have been reported as either individual case reports or as a subgroup in a series of infratentorial arachnoid cysts (Table 1).1,3,4,8,11,13,14,16,17,19,21,22

The indications for surgery and the type of surgical treatment are still subjects of debate.13 Not all arachnoid cysts require surgical intervention. Conservative management with regular radiological control for asymptomatic patients will identify those patients with gradual cyst enlargement and/or manifestation of new symptoms that could need surgical treatment.20 Consequently, in case of

FIG. 1. Intraoperative image obtained in a child with a CPA arachnoid cyst. After dura opening and gentle cerebellar retraction, the posterior wall of the cyst formed by thickened arachnoid is visualized. Figure is available in color online only.

FIG. 2. Intraoperative image obtained in the same case as Fig. 1 after opening of the cyst and extensive removal of the cyst wall. The abducens nerve and the CN VII-VIII complex are entirely dissected and decompressed. Figure is available in color online only.

FIG. 3. Preoperative (left) and postoperative (right) T2-weighted MR images showing the right CPA arachnoid cyst (left) and the complete reexpansion of the cerebellum 6 months after surgery (right).
incidental discovery of a CPA arachnoid cyst in an asymptomatic individual, a “wait-and-see” strategy could be justified, but the patient’s family should be aware that any initial loss of hearing or evidence of cyst enlargement at neuroimaging should prompt immediate consideration of the possibility of surgical treatment, as long-standing acoustic deficits seem to not respond or respond only partially to the cyst excision. In fact, in patients with a long history of hearing loss, the family should be informed about the possibility that surgery may not restore normal acoustic function.

Indications for operative management include the presence of cyst growth, signs of neural compression, or hydrocephalus.13

Different surgical techniques have been reported for the treatment of intracranial arachnoid cysts: microsurgical approach with cyst fenestration, marsupialization into the subarachnoid space and/or resection of the cyst wall, stereotactic puncture, and cystoperitoneal shunt placement or treatment of the associated hydrocephalus. Endoscopic management is becoming used more often. For the CPA arachnoid cyst in particular, direct attack of the cyst through a lateral suboccipital approach was predominant (Table 1).5,20 Aspiration techniques using stereotactic or endoscopic puncture9,10 are described in some reports, although reclosure of the cyst wall7 or insufficient fenestration resulting in inadequate drainage into the subarachnoid space are complications, leading to recurrence and regrowth. Cystoperitoneal shunting, as advocated by Ciricillo et al.5 also carries a risk of recurrence due to the occlusion of the system.12,15 Moreover, because of the close relationship to many important neurovascular structures in this area, cyst shunting, puncture, or endoscopic procedures may be hazardous in these patients.

Encouraging results in treatment and avoidance of recurrence have been observed in our previous experience with adult patients using a microsurgical retrosigmoid approach with resection of the cyst wall.20

Hearing loss is a rare presenting symptom of patients with CPA arachnoid cysts, described only in 2 previous pediatric cases reported in the literature.14,19 Although the exact pathophysiological mechanism for hearing loss in patients with arachnoid cysts has not been well described, various hypotheses have been proposed. Jayarao et al.14 suggested that their patient was symptomatic as a result of the arachnoid cyst exerting mass effect on the CN VII-VIII complex. It has been suggested that long-term compression of the cranial nerves in the CPA may cause irreversible nerve damage, most likely due to vascular injury. Even though it is often stated that children could have a better chance of recovery than adults because of the plasticity of the child’s nervous system, the duration of the clinical history in symptomatic cases appears to be significant. Indeed, by analyzing the cases presented in the literature14,19 and taking into account our own experience, we noticed that the patients with a longer preoperative duration of deficit had a worse hearing outcome (Table 2).

Also, considering all the cases presented in the literature, we can observe different outcomes: 1) complete resolution of the hearing deficit, as in the 2 children (one described by Olaya et al.19 and one in the present series) operated on shortly after diagnosis, who demonstrated immediate postoperative improvement of hearing deficit; 2) significant improvement, as in the patient described by Jayarao et al.14 who showed improved hearing at the 3-year follow-up visit but not in the immediate postoperative evaluation; and 3) only mild improvement, as in the child who was referred to our institute and treated 2 years after onset of hearing deficit, due to delayed diagnosis.

Conclusions

Cerebellopontine angle arachnoid cysts are uncommon in the pediatric population. The indications for treatment are of paramount importance in cases of symptom progression. If a hearing deficit is present, rapid surgical treatment is recommended to increase the chances of recovery.

References


TABLE 2. Cases of CPA arachnoid cysts in which patients presented with hearing deficits

<table>
<thead>
<tr>
<th>Authors &amp; Year</th>
<th>Time to Treatment</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jayarao et al., 2009</td>
<td>14 mos*</td>
<td>Unchanged</td>
</tr>
<tr>
<td>Olaya et al., 2011</td>
<td>6 mos</td>
<td>Compl resol</td>
</tr>
<tr>
<td>Present Case 1</td>
<td>3 wks</td>
<td>Compl resol</td>
</tr>
<tr>
<td>Present Case 2</td>
<td>24 mos</td>
<td>Unchanged</td>
</tr>
</tbody>
</table>

FU = follow-up. * Not precisely stated in the publication.

Disclosures
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: Giordano, Gallieni, Di Rocco. Acquisition of data: Gallieni, M Samii. Analysis and interpretation of data: Giordano. Drafting the article: Giordano, Gallieni. Critically revising the article: Giordano, A Samii, Di Rocco, M Samii. Administrative/technical/material support: Giordano, M Samii. Study supervision: A Samii, Di Rocco, M Samii.

Correspondence
Mario Giordano, International Neuroscience Institute–Hannover, Rudolf Pichlmayr Str. 4, Hannover 30625, Germany. email: giordano.nch@gmail.com.