Intracranian arachnoid cysts in children (ACs)

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Abstract
Intracranian arachnoid cysts (ACs) represent an extremely common condition in pediatric pathology. With the development of CT Scan and especially MRI these cysts could be find more constantly.
ACs are congenital lesions with maximum frequency in middle cranial fossa, followed by suprasellar aria, ponto-cerebelar angle and cranial posterior fossa. These cysts are often incidentally uncovered during a routine neuroimaging investigations for cranio-cerebral trauma or other diseases.
The authors present a series of 317 cases in children with ACs over a period of 10 years.
The authors avocate over the MRI evaluation of Acs and refering to therapeutic approach it is recomended only in compresive forms with focal neurologica sings or seizures.
Are rewiewed therapeutical procedures as: microsurgical fenestration with cystwall excision, endoscopic approach, stereotaxic suction, cyst shunting by cysto-peritoneal procedures. A number of cases remain under observation the surgical treatment beeing unnecesary. The surgical treatment must be carfully chosen, there is non therapeutical priority. It remains that improved neuroendoscopic methods to improve operator prognosis in Acs.
Keywords: Intracranial arachnoid cyst, MRI, microsurgical fenestration, endoscopic approach, cyst shunting, increased intracranial pressure, seizures

Introduction
Intracranian arachnoid cysts(ACs) also known as leptomeningeal cysts, are congenital, benign, nonneoplastic, extraxial lesions. ACs arise during development from splitting of arachnoid membrane, and are unrelated to infection.
Bright (1831) describe the intraarahnoidian origin of these lesion as “serous cysts forming in connection with the arachnoid and apparently lying between its layers.”, Starkman et all (1958) proposed that the asociated temporal hypoplasia is secandary to cyst expansion and pressure on the temporal operculum. Also Robinson (1961) the primary source of the problem lay in a congenital failure of temporal lobe development.

According to Di Rocco et all(2010), Acs are developmental defect, that occurs within the first three months of gestational life, in the duplication or splitting of the arachnoid layers, and are related to abnormalities of CSF flow. This theory cover the whole development of intracranial arachnoidian cysts. Acs are
associated with other developmental abnormalities of the brain, such as heterotopias. Incidence of Acs in 5 per 1000 in autopsy series and represent 1% of all intracranial masses. The male: female rate is 4:1, Di Rocco et all. (2010). Multiple or bilateral arachnoid cysts are unusual, and familial occurrence has been reported in only a few cases.

**Location**

In 50% intracranian arachnoid cyst involve the Sylvian fissure/middle cerebral fossa. Rengachary&Watanabe (1980) (Table I). According to Gallasi et all. (1980) Sylvian Acs can be classified into three subgoups in connection with cysts dimensions and extensions.

Gallasi type I: small, biconvex, located in anterior temporal tip, no mass effect, communicates with subarachnoid space. (Figure 1).

In Gallasi type II: involves proximal and intermediate segments of Sylvian fissure, completely open insula gives rectangular shape, partial communication with subarachnoidian space (Figure 2).

In Gallasi type III: involves entire Sylvian fissure, midline shift, bony expansion, minimal communication with the subarahnoidian space and surgical treatment usually does not result in efficient expansion of brain (Figure 3).

**Figure 1** MRI-aspect of Gallasi I left temporal arachnoid cyst

**Figure 2** MRI – aspect of Gallasi II left temporo-insular arachnoid cyst

**Figure 3** MRI- aspect of Gallasi III left cerebral hemisfere arachnoid cyst with middle line shift
Other common locations are: CP angle, the quadrigeminal cistern, the retrocerebelar area and the sellar/suprasellar region. Less commonly ACs can develope within the interhemisféric fissure and cerebral convexity (Tabel I).

**Clinical findings**

ACs become symptomatic mainly during childhood and adolescence, depending on the location of the cyst not to it’s dimension. Asymmetrical macrocranian or a focal bulging of the skull in the temporal region is the most common symptom, headaches, focal neurological symptoms, epilepsy and sings of increased intracraniel pressure.

In suprasellar ACs endocrine disfunctions (60% of cases), hydrocephaalus (40% of cases - probably due to compression of the third ventricle) and visual impairment are the most common presenting symptoms.

Natural evolution occuring without sings of: inflammation, trauma or hemorrhage. May be associated with other congenital anomalies (agenesis of the corpus callosum). Often do not expand and rarely may spontaneously regress or disappear.

**Terapeutical options**

Observation - Many authors recommend not treating arachnoid cysts that do not cause mass effect or symptoms, regardless of their size and location. Fatih E, Burkan B, Pinar O (2003)

Multimodal surgical treatment consisting of shunting the cyst into peritoneum, craniotomy (microsurgery) with fenestration and cystwall excision, endoscopic fenestration or stereotactic suction.Ciurea et all.( 2010)

Nowadays endoscopic fenestration represent the main therapeutical option in arachnoid cyst for decompression and restoration of CSF circulation.

Each procedures has advantages and disadvantages as described in table II.

**TABEL I**

**Localization of ACs(Rengachary&Watanabe 1981)**

<table>
<thead>
<tr>
<th>Location</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sylvian fissure</td>
<td>49%</td>
</tr>
<tr>
<td>CPA</td>
<td>11%</td>
</tr>
<tr>
<td>Supracollicular</td>
<td>10%</td>
</tr>
<tr>
<td>Pineal aria</td>
<td>9%</td>
</tr>
<tr>
<td>Sellar &amp; suprasellar</td>
<td>9%</td>
</tr>
<tr>
<td>Interhemispheric</td>
<td>5%</td>
</tr>
<tr>
<td>Cerebral convexity</td>
<td>4%</td>
</tr>
<tr>
<td>Clival</td>
<td>3%</td>
</tr>
</tbody>
</table>

**TABEL II**

**Surgical treatment options for arachnoid cysts (Keyvan Abtin and Marion L 2010)**

<table>
<thead>
<tr>
<th>PROCEDURE</th>
<th>ADVANTAGES</th>
<th>DISADVANTAGES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drainage by needle aspiration or bur hole evacuation</td>
<td>• siple&lt;br&gt;• quik</td>
<td>• high rate of recurrence of cyst and neurologic deficit</td>
</tr>
<tr>
<td>Craniotomy, excising cyst wall and fenestrating into basal cistern</td>
<td>• permits direct inspection of cyst&lt;br&gt;• loculated cysts treated more effectively&lt;br&gt;• avoid permanent shunt&lt;br&gt;• allows visualization of bridging vessels</td>
<td>• subsequent scarring may block fenestration allowing reaccumulation&lt;br&gt;• significant morbidity and mortality</td>
</tr>
</tbody>
</table>
Endoscopic cyst fenestration

- minimal invasive
- avoid permanent shunt
- no visualization of bridging vessels - hemorrhagic risk
- fenestration may close

Shunting of cyst into peritoneum

- low morbidity/mortality
- low rate of recurrence
- patient “shunt dependent” - risk of shunt infection

Patients and methods

The authors study 317 cases of arachnoid cysts, admitted in 1st neurosurgical clinic, pediatric neurosurgical department Bagdasar-Arseni Hospital between January 2002-January 2010 (8 years). Admission criteria was: age between 0 to 16 years old and patients diagnosed, treated and followed in 1st Neurosurgical clinic.

There were excluded all patients over 16 years old or treated in other neurosurgical services.

Results

Localisation of ACs: Sylvian fissure 172 cases (54%), CP angle 38 cases (12%), sellar and suprasellar region 32 cases (10%), pineal area 28 cases (9%), retrocerebelar 28 cases (9%) and interhemisferic 19 cases (6%). (table III)

<table>
<thead>
<tr>
<th>Location</th>
<th>Number of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sylvian fissure</td>
<td>172</td>
<td>54%</td>
</tr>
<tr>
<td>CPA</td>
<td>38</td>
<td>12%</td>
</tr>
<tr>
<td>Sellar &amp; suprasellar area</td>
<td>32</td>
<td>10%</td>
</tr>
<tr>
<td>Pineal area</td>
<td>28</td>
<td>9%</td>
</tr>
<tr>
<td>Retrocerebelar</td>
<td>28</td>
<td>9%</td>
</tr>
<tr>
<td>Interhemisferic</td>
<td>19</td>
<td>6%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>317</strong></td>
<td><strong>100%</strong></td>
</tr>
</tbody>
</table>

The most common clinical finding was the focal bulging of the skull 62% (196 cases), irritability 51% (162 cases), epilepsy 32% (101 cases) minor focal neurological symptoms 18% (57 cases), cranial nerves palsy 11% (35 cases) and increased intracranial pressure syndrome in 13% (41 cases) (table IV).

<table>
<thead>
<tr>
<th>Clinical findings in current study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Focal bulging of the skull</td>
</tr>
<tr>
<td>Irritability</td>
</tr>
<tr>
<td>Epilepsy</td>
</tr>
<tr>
<td>Focal neurological symptoms</td>
</tr>
<tr>
<td>Cranial nerves palsy</td>
</tr>
<tr>
<td>Increased ICP</td>
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</tbody>
</table>

Elected treatment was: (table V)

<table>
<thead>
<tr>
<th>Type of intervention</th>
<th>Number of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observation</td>
<td>40</td>
<td>12.6%</td>
</tr>
<tr>
<td>Unishunt cysto-peritoneal dr.</td>
<td>162</td>
<td>51.4%</td>
</tr>
<tr>
<td>Low pressure valv CPS</td>
<td>20</td>
<td>6.3%</td>
</tr>
<tr>
<td>Endoscopic procedure</td>
<td>31</td>
<td>9.7%</td>
</tr>
<tr>
<td>Mycosurgery</td>
<td>64</td>
<td>20.1%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>317</strong></td>
<td><strong>100%</strong></td>
</tr>
</tbody>
</table>
Case presentation I

A 16 years old girl presenting with migranal syndrome. The MRI investigation show a Gallasi tip I cyst of the left middle fossa. The elected treatment was observation with MRI reevaluation annually.

- observation in 40 de cases (12,6%)

Case presentation II

A 4 years old girl presenting with epileptic seizures. The preoperative investigation show a Gallasi tip II left middle fossa cyst. The elected treatment was a unishunt cysto-peritoneal shunt.

- unishunt cysto-peritoneal drainage 162 cases (51,4%) & low pressure valv cysto-peritoneal shunt in 20 cases (6,3%).

Case presentation III

A 3 years old boy presenting with seizures unresponsive to medical treatment. The MRI evaluation show a Gallasi III arachnoidian cyst of left cerebral hemisphere. The elected treatment was microsurgical approach with cyst wall excision and fenestration in 64 cases (20,1%).

- microsurgical approach with cystwall excision and fenestration in 64 cases (20,1%).

Case presentation IV

A 5 years old girl presenting with headaches. The MRI investigation show a Gallasi tip IV left occipital fossa cyst. The elected treatment was a microsurgical approach with cyst wall excision.

- microsurgical approach with cystwall excision in 64 cases (20,1%).

Case presentation V

A 7 years old girl presenting with headache and visual disturbances. The MRI investigation show a Gallasi tip V left occipital fossa cyst. The elected treatment was a microsurgical approach with cyst wall excision and fenestration.

- microsurgical approach with cystwall excision and fenestration in 64 cases (20,1%).
Microsurgical approach with cystwall excision and fenestration is recommended as an initial approach to avoid shunting. Shunt independence an important surgical goal. Keyvan Abtin & Marion L. Walker (2007) - endoscopic procedure in 31 cases (9,7%)

Case presentation IV

A 1 year old girl presenting with hypotonic syndrome and psychomotor retardation. The MRI examination show a supracollicular giant ACs, The elected treatment was endoscopic fenestration.
Discussion

Intracranial arachnoid cysts are mentioned throughout the specialised literature. The discovery of such cysts in almost all cases is done when is made an CT scan or MRI exam of the brain for other disease of nervous system. Once the intracranial arachnoid cyst is found, one has to establish very carefully the clinical findings, neuroimaging evaluation and after that we may proceed to a possible surgical treatment.

There are 2 histological types of ACs:

1. “simple arachnoid cysts”: arachnoid lining with cell that appear to be capable of active CSF secretion. Middle fossa cysts seem to be exclusively of this type.

2. cysts with more complex lining which may also contain neuroglia, ependyma and other tissue types. Mayr U, Aicher F, Bauer G et al (1982)

Many authors recommend observation and monitoring of these cysts on MRI. The rarity of expanding arachnoid cysts makes frequent serial neuroimaging unnecessary and cost ineffective. The recommended follow-up neuroimaging is at one year.

In case of a symptomatic cyst by neurologic focal deficit, epilepsy or raised ICP the best surgical solution to release pressure on the cerebral structure has to be found.

For a long time shunting methods were preferred. But with the appearance of shunt complication this methods where abandoned. Microneurosurgical approach of fenestration are very effective but they require large opening of basal cistern.

Nowadays as minimal invasive procedure the endoscopic procedures has become increasingly popular “the procedure of choice” and has been used to decrease the number of shunts. Hopf NJ, Perneczky A (1998)

As effective and safe and less invasive but great care is needed to avoid bleeding -ACs are associated with large bridging veins. F. di Rocco et al (2010)

Intraoperative ultrasound or frameless...
stereotaxy, especially with intravenous contrast enhanced CT or MR imaging guidance, is helpful in choosing the trajectory of the bridging veins. Main limitation is anatomical: due to the relation of the cyst with the basal cisterns and the temporal lobe displacement. C di Rocco et all (2010)

Conclusions
ACs are very frequent congenital intracranial malformation.
More than 80% of ACs are incidental findings being completely asymptomatic.
Treatment is recommended only in symptomatic ACs by focal neurological deficits, skull deformities, signs/symptoms related to increased ICP and seizures not responsive to medical treatment.
Current series of 317 cases constitute a uniform cohort because cases are diagnosed, treated and followed in a single pediatric neurosurgical service. Choice of treatment was performed very carefully to obtain the best clinical outcome and imaging properly, reducing the size of the cyst. Ciurea at all (2010)
All cases requiring follow predominantly by MRI to monitor the possible expansion of the cerebrale structures. In a significant number of cases the expected expansion did not occur because of the cyst membranes were not enough fenestrated into basal cisterns and CSF circulatia not restored properly. In these cases depending on clinical aspects the therapeutical process can be repeated, minimally invasive by endoscopy.
Basically we can say that there is no “Best Treatment” in ACs, and each case must be analyzed separately. Ciurea et all (2010)

Abbreviations:
ACs - intracranial arachnoid cysts
CSF - cerebro spinal fluid
CPA - cerebro-pontine angle
CPD - cisto-peritoneal drenage
MRI - magnetic resonance imaging
CT - computer tomography
ICP - intracranian pressure

References