Arachnoid Cyst with a Non-traumatic Acute Subdural Hematoma in an Eleven-year-old Patient

Cisto aracnoide associado a hematoma subdural agudo não traumático em um paciente de onze anos de idade

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Abstract
A non-traumatic intra-cystic hemorrhage in an arachnoid cyst is a rare event, with few cases reported in the literature. We present a case of an eleven-year-old boy patient, which presented a spontaneous acute subdural hematoma and intra-cystic hemorrhage after a strong headache episode. The results were evidenced by a computed tomography (CT) scan and surgical findings. We perform a brief literature review on the arachnoid cyst and its suggested treatments. In our case, the patient underwent a surgical treatment with a complete resolution of the case.

Keywords
► neurosurgery
► arachnoid cysts
► acute subdural hematoma

Resumo
Sabe-se que uma hemorragia intracística não traumática em um cisto aracnoide é um acontecimento raro, com poucos casos relatados na literatura. Nós apresentamos o caso de um menino de onze anos de idade, o qual apresentou espontaneamente um hematoma subdural agudo acompanhado de hemorragia intracística após um forte episódio de dor de cabeça. Os resultados foram evidenciados através de uma tomografia computadorizada e dos achados cirúrgicos. Foi feita uma breve revisão da literatura sobre cisto aracnoide e seus respectivos tratamentos sugeridos. Em nosso caso, o paciente foi submetido a um tratamento cirúrgico, com resolução completa do caso.

Palavras-chave
► neurocirurgia
► cisto aracnoide
► hematoma subdural agudo

Introduction
The arachnoid cyst is presumably a congenital malformation and represents only 1% of all intracranial lesions. Its rupture and bleeding are rare, usually occurring in young adults and associated with trauma. Risk of hemorrhage does not exceed 0.04%/year. We report a case of an eleven-year-old boy with a previously asymptomatic arachnoid cyst that suddenly presented a non-traumatic acute subdural hematoma and intra-cystic hemorrhage.

Case
An eleven-year-old boy had a sudden episode of headache of moderate intensity accompanied by nausea. The child denied any history of chronic headache before this episode. The symptoms began appearing at a monthly frequency. Every time, the boy’s parents would medicate symptoms with oral analgesic, after which he evolved with improvement. After five months, the headache episodes became weekly and more

intense associated with nausea and vomiting, varying in location. Oral analgesics were no longer effective to solve the pain. There were no other associated symptoms. The parents decided to take the patient to the emergency service, where he had a cranial CT scan. This one showed an arachnoid cyst in the middle fossa of the left cerebral hemisphere, without signs of complications (►Fig. 1). The local neurologist evaluated him and initiated a conservative treatment of the headache. After about a month with worsening symptoms, the patient returned to the emergency service, where he had a new cranial CT scan that showed an acute subdural hematoma (►Fig. 2). He was referred to the neurosurgeon local service, where there was a decision made to hospitalize the patient and provide surgical treatment.

The surgical technique was decompressive craniotomy with a question-mark incision and a whole hematoma evacuation. There was intracystic hemorrhage observed during the surgery. The patient remained in the pediatric intensive care unit due to postoperative support, showing gradual improvement. The patient received a medical discharge after seven days. After 30 days, he had a new CT. This last CT scan showed no bleeding. Based on the CT findings and the absence of pathological signs and symptoms, we concluded there had been a complete resolution of the condition (►Figs. 3–4).

Discussion

Intracranial arachnoid cysts are relatively common findings in neuroimaging studies among the pediatric population, primarily in boys. The most accepted hypothesis states that the arachnoid cyst is a congenital malformation.

The arachnoid cyst corresponds to 1% of all non-traumatic intracranial lesions. The most common location is the middle cranial fossa. It is presumably formed by a division of the arachnoid membrane caused by an increase in cerebrospinal fluid (CSF) pressure. Other hypotheses are: 1-dysgenesis embryo in the formation of the arachnoid, secondary to a primary disorder of the mesenchyme adjacent to the neural tube; 2-agenesis, located atrophy or hypoplasia of the brain resulting in the secondary expansion of the CSF space; 3-located disorder secondary to infectious/inflammatory process, traumatic or hemorrhagic. Most imaging studies findings are incidental due to the asymptomatic clinic nature of the arachnoid cysts. The risk of bleeding does not exceed 0.04% per annum. When bleeding does occur, most cases are associated with head injury. In this case, there was no report of injury, which makes this case even rarer. The most plausible hypothesis was spontaneous bleeding. According to Patel et al, the source of the bleeding was in the communicating vessels between the dura and the outer face of the cystic membrane. Bleeding may occur not only in the cyst, but in the subdural and epidural extra space as well. However,
intra-cystic hemorrhage is an extremely rare complication, which still remains with its pathogenesis unknown. This complication was visibly present in our patient. In a case-control study, we observed that the larger the cyst size, especially if the cysts have a maximum diameter larger than 5 cm, the greater the chance of disruption and associated complications, such as hygroma, subdural hematoma, and intra-cystic bleeding. Moreover, signs of intracranial hypertension, seizures, macrocrania, and delayed development in the last thirty days must be considered warning signs of cyst rupture.

Based on the literature reviewed, treatment is essentially surgical, although largely contingent on the patient’s symptoms. Surgery is indicated when there are important symptoms and/or complications of the disease. In asymptomatic patients, there is no consensus in the literature for the surgical treatment.

We found that patients with symptomatic arachnoid cysts can safely undergo a key-hole craniotomy and drainage procedure with reported success.

**Conclusion**

Arachnoid cysts are usually incidental findings in neuroimaging and more common in pediatric patients, appearing asymptomatic in most cases. Its pathophysiology is currently not well explained. The diagnosis is well established by a head CT without contrast or MRI. The literature also suggests that vascular imaging tests should be requested for patients with arachnoid cyst who presented headache.

The most common symptom in this case is the headache. Treatment is primarily surgical in symptomatic cases and/or when complications are present. Patients with arachnoid cyst with presence of warning signs should be investigated for rupture and bleeding of the lesion.

**Complementary Information**

We conducted this study at the institution Santa Casa de Misericórdia de Araçatuba, in the town of Araçatuba, in the state of São Paulo, Brazil.

**References**