Glial Cyst inside the Cerebellar Parenchyma: Case Report and Literature Review

Cisto glial no parênquima cerebelar: relato de caso e revisão da literatura

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Abstract
Cystic lesions inside the brain parenchyma are a common pathological finding in the investigation of patients with suspicion of cerebral tumor. Histological diagnosis is important to guide the patient's treatment and follow-up, and to determine prognosis. Among patients diagnosed with cerebral cysts, most are located in the parenchyma above the tentorium. The authors describe the case of a patient who had been suffering from dizziness and balance disturbance for 4 months; the investigation identified a cyst inside the cerebellar right hemisphere. A surgical procedure was performed, and the biopsy microscopic analysis diagnosis was glial cyst.

Keywords
► cerebellar cyst
► cysts inside the parenchyma
► glial cyst

Resumo
Patologias intraparenquimatosas de características císticas são comumente identificadas em pacientes que estão sob investigação de neoplasias cranianas. Lesões císticas são mais prevalentes acima da tenda cerebelar, e o diagnóstico histológico é fundamental para determinar o tratamento, seguimento e prognóstico do paciente. Os autores relatam o caso de um paciente com sintomas de vertigem e alteração no equilíbrio dinâmico de 4 meses de evolução, cuja investigação diagnosticou cisto intraparenquimatoso no hemisfério cerebelar direito. Procedeu-se com intervenção cirúrgica cujo diagnóstico histopatológico foi compatível com Cisto Glial.

Introduction

Case Report
A 63 year-old retired man was admitted to the neurosurgical department of Hospital Cristo Redentor (located in the city of Porto Alegre, Brazil) with a history of whole brain headache associated with vertigo and loss of balance for the four months previous to the medical evaluation. The patient had no history of head trauma or oncologic records. The neurological exam did not evidence sensitive or motor deficits, no deep or superficial reflex asymmetry, and cranial nerve tests did not evidence any impairment. Muscular tonus and tropism were normal. Cerebellar tests did not show any alteration. Although the patient did not stand still when he...
was asked to stand up and close his eyes, he did not fall. Dynamic balance was deviated to the left, a sign that was clearer when the patient tried to walk with one foot right in front of the other following a straight line drawn on the floor. The patient had no vertigo associated with fast passive head rotation, but his thought and speech was slow.

A computed tomography (CT) scan was performed, and it showed a hypodense and cystic lesion in the right cerebellar hemisphere, with no apparent solid tissue around or inside its borders. To better analyze the lesion and try to limit diagnoses options, full magnetic resonance imaging (MRI) was performed. The exam showed a cystic intracerebellar lesion, with a hypointense sign on the T1 series (►Fig. 1) and a hyperintense sign on the T2 series (►Fig. 2), but without contrast sign after a gadolinium infusion (►Fig. 3). The MRI also evidenced that the lesion did not have contact with the subarachnoid space or with the fourth ventricle. No artery or venous blood vessels were related to the cyst, and there was no solid tumor or apparent capsule surrounding it. Although the cyst did not have communication with the cisternal or

![Fig. 1](image1.png)  
**Fig. 1** Magnetic Resonance in the T1 weight series on sagittal view showing a cyst inside the cerebellar parenchyma.

![Fig. 2](image2.png)  
**Fig. 2** Magnetic resonance in the T2 weight series on coronal (A) and axial (B) views. Both images show a cystic lesion with intensity similar to the cerebrospinal fluid’s. A fourth ventricle deviation to the left side is also shown in both images.

![Fig. 3](image3.png)  
**Fig. 3** Magnetic resonance images in axial (A) and sagittal (B) views in the T1 series after a gadolinium infusion. In both images there is no contrast enhancement inside the cyst or its borders.
ventricular systems, the intensity of its content was similar to the liquor’s. The fourth ventricle was compressed and deviated to the left side. However, the patient had no clinical or radiological criteria for hydrocephalus.

The patient was then submitted to surgery to analyze the cyst’s content and pathological diagnosis. A lateral suboccipital access was performed, exposing the cerebellar right hemisphere. Once the surgical site was reached, a tan-yellow fluid was aspirated, without signs of recent or old hemorhages. A macroscopic inspection showed a regular parenchyma, and this impression was maintained during the microscopic view. The cystic walls were opaque with regular cerebellar tissue appearance, and no solid tissue was found during inspection. Neurosurgeons decided to perform random biopsies of the cyst wall, drain its fluid content, and send it to pathology and microbiology analyses.

The patient’s surgical recovery was excellent: he had no more headaches, could walk by himself without balance disturbance, and his speed of thought and speech improved. A CT scan after surgery showed a satisfactory cyst reduction (Fig. 4).

Laboratory results evidenced a basic fluid (pH value of 8.56) with lactate of 2.8mg/dL, elevated protein concentration (2000 mg/dL) and normal glucose value (60mg/dL). Microbiological investigation did not found any bacteria, fungus or bacillus tuberculosis on cultural exams. Cytological analyses demonstrated a liquid with rare epithelioid cells with no atypical nuclear and cellular structures. The histopathological microscopic structure evidenced three layers of regular glial cells surrounding the cyst, with a diagnosis of glial cyst (Fig. 5).

The patient was released from the hospital one week after surgery, without neurological impairments, a clinical status that remained the same during follow-up. However, a CT scan performed three months after surgery showed a reorganization, in smaller proportions, of a cystic lesion in the same anatomical place (Fig. 6). Since the patient was asymptomatic and the histological diagnosis of a benign pathology was known, the authors decided to follow this patient with CT scans and neurological evaluations every six months.

Discussion

Posterior fossa cysts located inside the parenchyma can be classified into neoplasm or non-neoplasm types. Pilocytic
astrocytomas, cystic teratomas, hemangioblastomas, metastases and ependymomas should be included in the differential list of the former type. The differential diagnosis of non-neoplasm posterior fossa cystic lesions includes many entities, such as infectious pathologies (toxoplasma abscesses, neurocysticercosis or hydatid cysts), arachnoid cyst, colloid, epidermoid or dermoid cysts. A least common differential diagnosis is the glial cyst, a very rare entity encountered in adults or older patients.

An article published in 1982 found 14 cases described in the medical literature from 1926 and 1981. A recent study conducted by Sundaram et al in 2001 analyzed 145 patients with diagnoses of cyst in the central nervous system across 12 years. Among these 145 cases, 2 (1.37%) were glial cysts, with diagnoses of cyst in the central nervous system across 12 years. Among these 145 cases, 2 (1.37%) were glial cysts, with diagnoses of cyst in the central nervous system across 12 years. Amically, no signs of tumor nodules are found, and the surgeon must perform random biopsies of the cystic wall for pathologic analyses. Once the pathological diagnosis of glial cyst is confirmed, surgical excision (if not performed during the first surgery) must be performed and is curative.

The proximity of the cyst to the fourth ventricle in the two cases reported by Vaquero et al made the authors hypothesize that the cyst originated from an ependymal diverticulum during intrauterine development, an idea first proposed by Szigethy. Some authors believe that they are probably formed by budding from the ventricular system, like an ependymal cyst with subsequent loss of ependymal cells by stretching or pressure effects. However, other authors accept as more plausible the hypothesis that a simple glial cyst represents an aborted or degenerated cerebellar astrocytoma, especially because of the fact that the reported glial cysts of the cerebellum are usually diagnosed during the fourth and fifth decades of life, allowing the assumption of degenerative changes in a previous undiagnosed cerebellar astrocytoma.

Most cases are asymptomatic, and the diagnosis occurs incidentally during imaging exams. However, they may become symptomatic because of pressure, rupture, or a secondary inflammation. Among the symptomatic cases, symptoms of increased intracranial pressure secondary to the cyst itself or obstruction of the fourth ventricle are the most common clinical manifestations. Both cases described by Vaquero et al presented a clinical evolution of a slowly growing expansive lesion. In our case, the cystic growth compressed the fourth ventricle, which impaired cerebrospinal fluid flow, causing the symptoms of headache and bradipsychism.

The first CT findings of cerebellar glial cysts were published in 1981. The cyst usually is homogeneously hypodense, with a Hounsfield count similar to the cerebrospinal fluid’s. An important characteristic is that glial cysts are not vascularized, and, therefore, do not enhance after contrast injection. On MRI, the cystic lesions have a signal characteristic of cerebrospinal fluid, a pattern showed during the radiological investigation of the case here described.

Despite the well-established characteristics on image exams, neuroradiological analysis without histological verification does not predict the benign nature of the lesion. A histopathologic study is important for diagnosis, to guide therapy and to establish patient prognosis. Therefore, special attention must be paid to obtaining adequate surgical specimens in order to distinguish a glial cyst from a more aggressive pathology, such as low-grade neoplasm.

On macroscopic inspection, the cyst is tan-yellow or opaque, and has a smooth capsule that may be up to 2 mm thick. The fluid content is clear yellow or yellow-white (from elevated protein content), and is not xanthochromic. These cysts may be unilocular or multilocular, with a central or lateral location, typically confined to the white matter, but they may also affect gray matter.

Microscopically, the appearance of the cyst wall is characteristic, revealing the cyst surrounded by bands of fibrous glial tissue. The wall is composed of three layers of neurologic structure with no epithelial lining. The outer layer consists of a fibrous capsule. The fibroblasts are sometimes arranged in a thin layer, but are more commonly densely packed. The inner layer is formed by a thin lining of glial cells. No mural nodules, tumor cells, or evidence of old or recent hemorrhages can be detected.

During surgery, once the cyst is reached, the surgeon must carefully look around the cystic cavity to try to identify a nodular or solid area to guide surgical biopsy. Characteristically, no signs of tumor nodules are found, and the surgeon must perform random biopsies of the cystic wall for pathological analyses. Once the pathological diagnosis of glial cyst is confirmed, surgical excision (if not performed during the first surgery) must be performed and is curative.
The radiological recognition of these cysts and diagnosis confirmation by pathology is essential for prognosis, which is extremely good once surgical evacuation of the cyst is performed.6

Conclusion

In a late middle-aged patient with appropriate cerebellar signs, a neuroradiological demonstration of a well-defined, non-enhancing cystic lesion on CT and MRI scans makes the diagnosis of a simple cerebellar cyst probable. However, the neurosurgeon must always exclude tumor and infectious pathologies, once they have a different prognosis and specific treatments. Besides, any tumoral growth in the posterior fossae may obstruct or divert the fourth ventricle, causing hydrocephalus and increasing intracranial pressure. In these cases, surgical resection is indicated.

References
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