Case Report

Xanthogranulomatous colloid cyst: Radiologic–pathologic correlation and diagnostic difficulties

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ABSTRACT

Background: Despite colloid cyst in the third ventricle is a very usual cause of hydrocephalus, its xanthogranulomatous variant is rare. The most important differential diagnosis is the third ventricular craniopharyngioma. To the best of the authors’ knowledge, there have been few cases of xanthogranulomatous variant colloid cysts reported in the English literature.

Case Description: A 77-year-old white woman presented with headaches, memory loss, and abnormal gait for the past 4 months. Magnetic resonance imaging revealed a solid cystic lesion measuring 3.0 cm×2.8 cm×2.9 cm located inside the anterior portion of the third ventricle causing obstructive hydrocephalus. The posterior portion of the lesion was predominantly solid and hypointense on T2 and T1, with areas of post-contrast enhancement, and the anterior portion was predominantly cystic with both hyper- and hypointense areas on T1 and T2, with no suppression on fluid-attenuated inversion recovery and no restriction to diffusion. The patient underwent a left frontal craniotomy with pterional approach, and the lesion was removed microsurgically.

Conclusion: Xanthogranulomatous reaction is rarely described in colloid cysts, which happens as a response to desquamation of epithelial lining, subsequent lipid accumulation, and as tissue inflammatory response to intracystic hemorrhage. Microsurgical resection is the treatment of choice. As compared to the plain colloid cyst, these lesions are difficult to fully excise as the inflammatory reaction to the xanthomatous material leads to adhesions to adjacent structures; therefore, the aspiration of cystic contents without spillage is advisable to achieve maximal resection of cyst walls.

Keywords: Mimics, Neuroimaging, Neuroradiology, Neurosurgery, Pathology

INTRODUCTION

Although colloid cysts are the most common benign neoplasm of the anterior third ventricle, its xanthogranulomatous transformation has scarcely been reported in the literature and may impact adequate patient management. Here, we report a rare case of atypical colloid cyst treated with neurosurgery and review the most significant imaging and surgical findings.
CASE REPORT

History

A 77-year-old white woman presented to the emergency room with complaints of headaches, memory loss, and abnormal gait for the past 4 months. She had a 10-year history of a controlled type 2 diabetes mellitus and hypertension. At neurological examination, she was awake, partially oriented, with pupils equal round and reactive to light, without papilledema, perceived strength, eumetric intention tremor, no frontal release signs, and ataxia of gait.

Radiographic features

A magnetic resonance imaging (MRI) revealed a hyperintense T1WI solid cystic lesion measuring 3.0 cm×2.8 cm×2.9 cm located inside the anterior portion of the third ventricle causing obstructive hydrocephalus. The posterior portion of the lesion was predominantly solid and hypointense on T2WI and on gradient echo (GRE) T2WI, and the anterior portion was predominantly cystic with both hyper- and hypointense areas on T1WI and T2WI, with no suppression on fluid-attenuated inversion recovery (FLAIR) and no restriction to diffusion. The solid portion was hypointense on T1WI and demonstrated heterogeneous areas of enhancement after gadolinium contrast agent administration; there were also some peripheral sparse areas of enhancement in the cystic portion of the lesion. These heterogeneous peripheral enhancements and hypointensity on GRE TW2I are consistent with either hemosiderin or calcifications, raising concern for an atypical colloid cyst or a cystic craniopharyngioma [Figure 1].

Surgical management

A left frontal craniotomy with pterional approach and a transfrontal route using endoscopic visualization and stereotactic guidance were performed, and the intraventricular lesion was removed microsurgically. The solid part of the lesion was excised, and most of the capsules were removed except for the part densely adherent to the floor of the third ventricle. About 3–4 ml of "machine-like oil" fluid was also drained.

In the intraoperative view, the lesion was solid cystic located in the third ventricle near the foramen of Monro. The specimen received and fixed in 10% buffered formalin, displayed a nodular appearance, measured 2.5 cm×2.0 cm×2.0 cm, and weighed 7.0 g. It was surrounded by a delicate irregular membrane, with a yellowish color and friable areas. The solid part was firm, grayish-black in color, and avascular [Figure 2].

Figure 1: Axial T1 (a) and T2 gradient echo (b); coronal T2WI (c) and sagittal post-Gadolinium T1WI (d). Heterogeneous solid-cystic mass obstructing the third ventricle, located at the foramen of Monro (blue arrow), at the anterosuperior aspect of the third ventricle, causing a deviation of the midline (white arrow) and obstructive hydrocephalus. The anterior portion of the lesion is predominantly cystic with a moderate hyperintense signal (yellow arrows). The posterior portion is predominantly solid (red arrow), has hypointense areas suggestive of either calcifications or a hemorrhagic component (green arrow), and shows heterogeneous enhancement (pink arrow).

Figure 2: Intraoperative images (a and b) microsurgical removal of the colloid cyst's capsule (blue arrow) using endoscopic visualization and resection of the colloid cyst (b). Gross specimen (c) displaying nodular appearance surrounded by a delicate irregular membrane, with a yellowish color (yellow arrows) and friable areas (inside white ellipses), size: 2.5 cm×2.0×2.0cm and weigh: 7.0 g.

Pathology

In the intraoperative histological examination, a diagnosis of craniopharyngioma was made, but the definitive histopathological examination of the excised specimen
revealed a colloid cyst lined by a single layer of cuboidal, partially flattened nonciliated and ciliated epithelial cells. There was a high prevalence of macrophages and epithelioid cells, many of which included refractile material and hemosiderin. In addition, cholesterol clefts and lymphocytes were seen in focal sets. Polymorphonuclear leukocytes were diffusely spread throughout the granuloma [Figure 3]. The final diagnosis of a xanthogranulomatous colloid cyst with hemosiderosis was made.

The immediate post-operative MRI demonstrated an almost complete reduction of the cystic lesion, with some sparse areas of contrast agent enhancement, and normalization in the lateral ventricle size [Figure 4]. Clinically, the patient evolved with the gradual improvement of the condition; she was discharged and followed-up conservatively with serial neuroimaging.

**DISCUSSION**

Colloid cysts account for 0.5%–1% of all intracranial tumors, with a yearly incidence of 3.2 cases per 1 million people; however, they are the most common benign neoplasm of the anterior third ventricle. The peak incidence is around the forties, and there is no gender prevalence\(^1,8\). They usually occur at the level of the foramen of Monro, and patients usually present with increasing intracranial pressure from cyst expansion and blockage of CSF flow. This benign neoplasia clinically manifests as an intermittent headache, characterized by short-duration attacks and relieved when sleeping (differentiating from brain-occupying lesions); nausea and vomiting can accompany headache episodes. In addition, sudden episodes of loss of consciousness and gait disturbances can be present, and, in a minority of patients, memory disturbance due to forniceal compression, seizures, and recurrent aseptic meningitis are described.\(^4,10,11\)

The congenital nature of colloid cyst is usually accepted, with origins ranging from primitive neurectoderm, choroid plexus, and ependyma. At gross pathology, the colloid cysts are usually ovoid well-delineated cysts with a thick gelatinous center with variable viscosity with rare evidence of hemorrhage. At microscopy, these tumors are usually described as being cystic and lined by a single or multilayered ciliated, cuboidal epithelium. A xanthogranulomatous reaction is rarely described in these cysts, which happens as a response to desquamation of epithelial lining and subsequent lipid accumulation as well as tissue inflammatory response to intracystic hemorrhage. Xanthogranulomas are seen as cholesterol crystal clefts, foreign-body giant cells, foamy cells, and inflammatory infiltrate of varying degrees located in the wall of colloid cysts.\(^2,5-7\)

Computed tomography (CT) studies showed that colloid cysts are hyperdense in two-thirds of cases and iso- to hypodense in one-third of cases, with density depending on the cholesterol content of the lesion. It usually does not enhance on contrast except for occasional rim enhancement.

On MRI, the signal varies according to the contents of the cyst, the characteristic appearance of being hyperintense on T1W images reflects the amount of cholesterol content, and the iso- to hypointensity on the T2W images is given by the amount of dense or proteinaceous fluid. There may be a peripheral (rim) gadolinium enhancement in a minority of
cases. These colloid cysts do not normally suppress on FLAIR studies and show no restriction on diffusion-weighted imaging.

Both CT and MRI studies may not be able to distinguish between colloid cysts and its xanthogranulomatous transformation; however, atypical features should alert for this differential diagnosis. The presence of blood within the cyst can trigger to xanthogranulomatous inflammatory response, which is seen as heterogeneous signal intensity on MRI; in these cases, signal characteristics are variable and depend on the mixture of lipid, fluid, and blood products.\[3,9\]

The presence of an atypical solid-cystic lesion in the third ventricle near the foramen of Monro, as in our case, can mimic a suprasellar cystic craniopharyngioma, especially the papillary subtype (considering patient's age), although such neoplasms are rarely presented in the third ventricle and more often have calcifications and gadolinium enhancement. Xanthogranulomas are often present in the atria of lateral ventricles and rarely can manifest in the third ventricle. They should be taken into consideration in the differential diagnosis of xanthogranulomatous colloid cysts, with iso- to a hyperintense signal on T1W images and variable gadolinium enhancement patterns. However, they have similar features and may be indistinguishable on imaging. In addition, xanthogranulomatous changes within colloid cysts or mixed tumors consisting of both of colloid cysts and xanthogranuloma have been described.

Other lesions, such as subependymoma and choroid plexus papilloma, can also be located in the foramen of Monro; however, the first is usually located at the 4th ventricle and demonstrates T2/FLAIR hyperintensity, and the second typically manifests in early childhood with an enhancing lobulated (cauliflower-like) mass in the atrium of the lateral ventricle.

Surgery is the treatment of choice, which can be accomplished either by endoscopic or microscopic techniques according to the expertise of the surgeon. As compared to the plain colloid cyst, these lesions are difficult to fully excise as the inflammatory reaction to the xanthomatous material leads to adhesions to adjacent structures, such as the walls of the ventricles.\[9\] Therefore, the aspiration of cystic contents without spillage is advisable to achieve a safe maximal resection of cyst walls.

The prognosis for xanthogranulomatous colloid cyst is usually good, depending on the amount of resected tissue. The most common complications, such as aseptic meningitis and vasospasm, arise post-operatively and are due to spillage of the cyst contents.\[1,10\] In the present case report, the patient evolved with the gradual improvement of the condition. She was discharged and followed-up conservatively with serial neuroimaging.

CONCLUSION

We reported a rare case of xanthogranulomatous colloid cyst of the third ventricle. Although computed tomography and magnetic resonance imaging may not be able to provide high specificity for this diagnosis, the presence of atypical neuroimaging features should raise concern, since this entity impose neurosurgery difficulties and its prognosis depend on the amount of resected tissue.

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Declaration of patient consent

The authors certify that they have obtained all appropriate patient consent forms. In the form, the patient has given her consent for her images and other clinical information to be reported in the journal. The patient understands that her name and initials will not be published, and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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Conflicts of interest

There are no conflicts of interest.

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