Intraventricular arachnoid cyst of lateral ventricle in an elderly patient

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Abstract
Arachnoid cysts are usually located in relation to the arachnoid cisterns. Intraventricular location is rare and its embryological emergence in this site is controversial. We report a large intra-ventricular cyst in a 61-year-old female who presented with decreased vision, headache and right hemiparesis. MRI was suggestive of cystic lesion in the lateral ventricle and was excised completely through a craniotomy.

Introduction
Arachnoid cyst comprises about 1% of intracranial space occupying lesions and are located in relation to arachnoid cisterns, most common locations being sylvian, chiasmatic, suprasellar, quadrigeminal and cerebellopontine cisterns. Symptomatic presentation of an intraventricular arachnoid cyst is very rare especially in the seventh decade of life and hence it being reported.

Case Report
A 61-year-old lady presented with decreased vision and headache of about one-year duration and weakness in the right side of the body for the preceding two and half months.

On examination: She was conscious; visual acuity was limited to perception of light in the right eye and finger counting at a distance of 2ft in left eye. There was increased tone in all limbs and right hemiparesis with a power grade III. A retinal hole had been detected in the right eye in another hospital.

Investigations: routine blood investigations, urinalysis, kidney function, liver function, ECG were reported to be normal. MRI of the brain revealed a large lesion, hypo-intense on T1WI and hyper-intense on T2WI suggestive of cystic mass in the lateral ventricle (Fig 1).
Craniotomy revealed large ventricle containing well defined thin walled cyst which could be easily separated from the ependyma. It was excised in-toto after coagulation of its attachment to the vessels. Histopathology of the cyst wall confirmed it to be arachnoid cyst.

Postoperatively her vision improved to finger counting at 6ft and regained full power in the right sided limbs. Her vision continued to improve over the period of observation to 6/16 and 6/24 which could be improved to 6/9 with glasses.

**DISCUSSION**

Most of the arachnoid cysts are asymptomatic and are discovered incidentally on imaging for other indications \(^1\). When they are symptomatic, the presenting features most often include headache, vomiting and seizures. Park et al collected 22 cases of arachnoid cyst of lateral ventricle from the literature and added one of their own \(^2\). Of these, 15

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**Figure 1.** Showing imaging characteristics of the intraventricular arachnoid cyst. On contrast CT (1a), it shows no enhancement of the cyst wall. The cyst is hypointense on T1 (1b), hyperintense on T2 (1d) with no wall enhancement on T1 contrast (1c).
were adults and 8 were children. Headache was the most common presenting feature in this series. Wong et al in 1993 reported a case which presented with positional psychosis due to intermittent blockage of temporal horn that occurred when the patient had been recumbent for 2-3 hours (3). Focal neurological deficits like hemiparesis as seen in our patient are rare at presentation. These lesions are seen usually in children or in the 4th decade of life. Recently, relatively large series of pediatric patients treated endoscopically have been published (4). However; our patient seems to have been the oldest in the 7th decade of life. What process had altered the status quo between the cyst and the patient is debatable as she had been apparently living with it for almost all of her life.

There are primarily two theories about the embryological origins of the arachnoid cysts. The first is the ‘arachnoid splitting theory’ (5). As the name suggests, it proposes that these cysts arise as a result of congenital splitting of the arachnoid layer wherein, the CSF accumulates gradually. The other theory is in specific context of middle cranial fossa arachnoid cysts and suggests that these arise as a result of temporal lobe agenesis as the primary event. However, it is named as ‘subarachnoid cyst theory’ (6).

There are various mechanism proposed for expansion of the arachnoid cysts. These include, but are not limited to, fluid secretion by cyst wall, presence of osmotic gradient and ball valve mechanism (7).

As mentioned previously, arachnoid cysts are purported to arise by a congenital splitting of the arachnoid membrane and subsequent accumulation of CSF in this ‘potential space’. However, normally, there is no arachnoid tissue in the ventricle. Thus, the origin of an arachnoid cyst in intraventricular location is controversial. According to Yeates and Enzmann, intraventricular arachnoid cysts arise from the vascular mesenchyme by invagination into adjacent brain, picking up an outer covering of adjacent glial tissue (8). Nakase et al. postulate that cyst arises from the arachnoid layer brought with vascular mesenchyme which it invaginates via the choroidal fissure (9). Usually, the cysts are located in the occipital horn and/or trigonal region and cause dilatation of the temporal horn and/ or occipital horn. In our patient, the cyst pathogenesis seems to mirror the mechanism proposed by Nakase et al. as it had an attachment to the choroidal fissure.

Imaging characteristics are those of any CSF containing cavity viz. hypo-intense on T1, hyper-intense on T2 and without any contrast enhancement. Symptomatic cysts require surgical intervention. The options available include endoscopic fenestration, cyst de-roofing, partial or complete removal and cysto-peritoneal shunt. Our patient was treated by complete removal of cyst wall through a craniotomy. Endoscopic fenestration was the safer and less invasive option. In Park’s series, there were various combinations and permutations of procedures applied for management of adult as well as pediatric intraventricular arachnoid cysts-open removal, open fenestration or partial cyst removal and endoscopic fenestration and shunt (2). Kurokawa et al recommended cyst resection rather than shunt because he found that the size of the ventricle was not reduced after the shunt procedure and the shunt malfunctioned due to wrapping of the shunt by the collapse cyst (10). Park et al had case of large arachnoid cyst of the lateral ventricle extending from the suprasellar cistern. The cyst wall could be easily separated from the ventricular ependymal after coagulation and transection of the attachments.

Cysto-peritoneal shunt or complete removal of the cyst wall are necessary to prevent recurrence, whereas cyst opening alone is insufficient. Our patient was treated by complete removal of the cyst wall through a craniotomy to avoid recurrence.

**Conclusions**

The embryological origin of intraventricular arachnoid cysts is controversial and the jury is still out. Our intraoperative observation during the cyst excision had shown the cyst to have an attachment to choroid fissure of the left lateral ventricle. This is in agreement with Nakase et al.’s propounded theory that intraventricular arachnoid cyst arises from the arachnoid layer brought with vascular mesenchyme which it invaginates via the choroidal fissure.

**References**