Case Report: Paired/Dual Colloid Cysts in the Third Ventricle - A Discussion of Embryological Origin

SM Jay* and MR MacFarlane

Department of Neurosurgery, Christchurch Hospital, Christchurch, New Zealand

*Corresponding author: SM Jay, Department of Neurosurgery, Christchurch Hospital, Christchurch, New Zealand, Tel: +6433641217, Fax: +6433641030, E-mail: sharonj2@cdhb.govt.nz

Abstract
A 31-year-old male presented with a 4-month history of rising intracranial pressure. Initial computed tomography (CT) and then magnetic resonance imaging (MRI) showed moderate dilatation of the lateral ventricles with periventricular signal change, an indistinct third ventricle with an ill-defined lesion in the anterosuperior part of the third ventricle. At time of operation two paired/dual colloid cysts were found lying side-by-side in the third ventricle and obstructing both foramina of Munro. This is the first described case of paired/dual colloid cysts in the third ventricle and re-ignites the debate as to the embryological origin of colloid cysts.

Keywords
Colloid cyst, Neuroepithelial cyst, Paraphysis, Third ventricle, Hydrocephalus, Neuroembryology

Clinical Details
A 31-year-old male presented to his local hospital with a 4-month history of recurrent mild headaches which became more severe in the week prior to admission, the severity of the headache reaching the point where he was unable to sleep and was associated with vomiting over the 24 hours before admission. Examination on admission showed he was fully alert and orientated (GCS 15/15) with no focal or abnormal neurological signs. Fundoscopy showed bilateral papilloedema with haemorrhages. An initial CT scan (Figure 1) showed moderate lateral ventricular dilatation with transependymal spread of cerebrospinal fluid (CSF) and a 2-3 mm hyperdense lesion in the anterosuperior aspect of the third ventricle - presumably a colloid cyst. The 4th ventricle was normal. An MRI scan (Figure 2) showed indistinct anatomy in the anterosuperior aspect of the 3rd ventricle and the appearances were not typical of a colloid cyst. Given the acute hydrocephalus and that there was an obstructive lesion in the anterosuperior aspect of the 3rd ventricle and that this was likely to be a colloid cyst we felt that direct surgery was undoubtedly indicated. The operation performed was a precoronal right frontal craniotomy and a transcortical approach via the right middle frontal gyrus and using microsurgical techniques to explore the anterosuperior aspect of the 3rd ventricle via the right foramen of Munro with the expectation of finding and removing a colloid cyst. A colloid cyst was found and removed, the thin wall being breached with removal, with the cyst containing glairy fluid. Once this cyst was removed a second cyst of similar size and appearance was seen lying to the left of the removed cyst and this second cyst was also removed, great care being taken so as to not breach the cyst wall so this cyst was kept intact. Essentially the cysts had been were sitting side by side. The wall of the first (right) cyst and the intact second (left) cyst were subjected to microscopic pathological examination and both lesions had the typical appearances of colloid cysts with ciliated cuboidal epithelium and the second (left) cyst was confirmed as being intact/unruptured. The patient had an uncomplicated postoperative course with a satisfactory post-operative MRI showing resolving hydrocephalus and restoration of normal 3rd ventricular anatomy. He was discharged from hospital neurologically intact on the 7th postoperative day. At outpatient follow up 3 months postoperatively he was headache free, neurologically intact, had normal memory functioning and was back at work.
Discussion

There have been three documented case reports of multiple (two) colloid cysts - the most recent by Dahdaleh, et al. [1] with a colloid cyst in each of the third and lateral ventricles, the second such case report of this kind [2]. Parkinson and Childe [3] postulated the mechanism for multiple colloid cysts, in their case both cysts in the fourth ventricle, was either originating from the ependymal lining within the fourth ventricle itself or a migration - where the cyst detached from the third ventricle and travelled through the aqueduct to attach in the 4th ventricle and increase in size to the point where they became symptomatic.

However, the authors present the first reported case of dual/paired colloid cysts in the 3rd ventricle and obstructing the foramena of Munro. This re-opens the debate into the embryological origin of these benign lesions. There are three major documented theories as to their pathogenesis.

1. Derivation from embryonic paraphysis or ventricular roof (neuroectoderm): The paraphysis first described by Salenka in 1890 as ‘a vesicle evaginating from the roof of the secondary forebrain or telencephalon and developing into a tubular structure [4]. The human paraphysis starts development from the telencephalic paraphyseal arch at 15-17 mm crown-rump length at approximately week 7 gestation. Kappers then found evidence of this between development stages 17-100 mm in 30 specimens and then postulated it degenerates at about the 100 mm stage (approx 3.5 months gestation). The U-shaped infolding of the diencephalic roof forms the velum interpositum [5] can undergo evagination into the space between the fornices or invagination into the third ventricle to form a colloid cyst- thus a colloid cyst is a midline structures by definition.

2. Developing from ependyma or choroid plexus: Kondziolka [6] used histochemical techniques to determine that colloid cyst epithelium is distinct, in terms of antigenic profile, from that of choroid plexus or ependyma, largely discounting this theory.

3. Ectopic respiratory/foregut epithelium: Lach, et al. [7] undertook immunohistochemical study of the colloid cyst epithelium from 21 colloid cysts and showed the immune phenotype of the epithelial lining to be similar to that of endoderm or ectoderm derived from respiratory or foregut origin. They compared this epithelium to the epithelium from Rathke’s cleft cysts and also follicular cysts of the pituitary and these were also similar in nature. They suggested a common histogenesis with these other neuroaxis cysts from the foregut-derived epithelia.

Conclusion

We believe our described case of bilateral/dual colloid cysts in the 3rd ventricle is the first such documented case report with colloid cysts found in such close physical proximity, each sitting side by side in the 3rd ventricle and each obstructing a foramen of Munro. Our case, combined with the previous reports of two more physically-separated colloid cysts, calls into question the embryological origin, certainly in relation to theory 1 above, that they are derived from the embryonic paraphysis or ventricular roof.

Conflict of Interest

The authors declare no conflict of interest. The authors alone are responsible for the content and writing of the paper.
References


