Endoscopic monoportal removal of a choroid plexus papilloma in the posterior third ventricle in a child

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Currently, only a few reports describe the minimally invasive removal of choroid plexus papillomas (CPPs) and, to the best of the authors’ knowledge, no reports detail the resection of such a papilloma through an endoscopic approach in infants. The authors here describe the endoscopic removal of a third ventricle CPP in a child. A 5-month-old male infant presented with progressive macrocephaly, vomiting, and convulsions. A lesion in the posterior third ventricle was detected on brain MRI. Because of the patient’s very young age, neuroendoscopy was used as the least invasive technique. The tumor was completely resected through a monoporal neuroendoscopic approach. Histologically, the tumor was classified as a WHO Grade I CPP. After surgery, the patient’s condition improved, with no complications during his recovery. Ten-month follow-up neuroimaging revealed no evidence of tumor recurrence or progressive hydrocephalus. In view of the successful neuroendoscopic excision of this posterior third ventricle CPP, the authors believe that this method seems promising in the treatment of young children with intraventricular lesions.


KEY WORDS choroid plexus papilloma; third ventricle; neuroendoscopy; monoportal removal; technique

choroid plexus papilloma (CPP) is a benign neoplasm that stems from cells of the choroid plexus and has a neuroectodermal genesis. Its annual incidence is 0.3 case per 1 million persons, and it constitutes 0.5%–0.6% of all intracranial tumors in all age groups. Few publications describe CPP removal in children via endoscopy either as an assisting tool or as an independent approach. Fully endoscopic removal of a third ventricle CPP has not been reported in the literature. In fact, third ventricle CPPs are rare, and their posterior location inside the third ventricle is even rarer. Here, we describe our experience in removing a posterior third ventricle CPP by adopting a fully endoscopic approach, and then we review the pertinent literature on the matter.

Case Report

History

A 5-month-old male infant was admitted to a local clinic when he developed generalized convulsive activity, anterior fontanelle bulging, and vomiting. Brain CT scanning revealed a mass in the third ventricle and internal hydrocephalus. His head circumference progressively increased. The boy regressed developmentally and was unable to roll over or hold his head up. He was transferred to the pediatric neurosurgical unit at the Federal Centre of Neurosurgery, Tyumen, several weeks later for further treatment. The period between the onset of symptoms and surgical intervention may have been used for diagnostic examination given the potential for slow progression of the disease. The delay may also be explained by geographical factors.

Upon admission to our institution, the boy remained unable to hold his head up or roll over. The patient had no complaints as regards the cardiovascular, respiratory, digestive, and urinary systems. His head circumference measured 46 cm, and the anterior fontanelle was taut. Neurological assessment revealed convergent strabismus, a decrease in muscle strength to +4 in the upper and lower extremities bilaterally, a decrease in bilateral upper and lower extremity muscle tone, hyperactive reflexes in the upper and lower extremities, and a Babinski sign bilaterally. Fun-
Duscopic examination was negative for congestion in the ocular fundus. Magnetic resonance imaging of the brain revealed an irregularly shaped mass, 2.12 × 1.61 × 1.45 cm in size, inside the third ventricle, which had enlarged to 2.21 cm. The mass had irregular, well-defined margins and was iso- to hypointense on T1-weighted imaging and mildly hyperintense on T2-weighted imaging. The mass occupied the posterior areas of the third ventricle, compressing the ventricular aqueduct and dislocating anterior aspects of the quadrigeminal plate into the cistern. Contrast injection revealed intense irregular enhancement of the mass. The lateral ventricles were enlarged asymmetrically, with the left larger than the right. Periventricular edema around the third and lateral ventricles was evident on T2-weighted and FLAIR imaging (Fig. 1).

Operation

The patient was selected to undergo monoportal endoscopic removal of the tumor, and the senior author performed an endoscopic ventriculocisternostomy (ETV) via the floor of the third ventricle. The patient was placed supine with his head immobilized inside a Doro clamp (PMI). We used endoscopic instruments for cranial surgeries: EndoWorld Neuro 12-E Lotta and Hopkins II 0° and 30° rigid endoscopes (both Karl Storz). Surgical entry into the anterior horn of the right lateral ventricle was performed parasagittally from the right frontal area via a bur hole. A neuroendoscope was then advanced inside the cranium. Tumor located in the third ventricle was visualized via the foramen of Monro (10 mm in diameter). Because MRI indicated that the tumor was protruding into the ventricular aqueduct and occluding it, we initially considered only a partial tumor resection. Given the occlusive nature of the hydrocephalus and in an effort to control its progression and avoid the need for a CSF diversion surgery in the future, we performed an ETV using a standard technique. Upon examination of the foramen of Monro and the third ventricle, we discovered that tumor tissue was intimately connected to the thalamostriate vein and the choroid plexus. After coagulation, the tumor tissue was sent for histological testing.

Working bimanually and using biopsy forceps, endoscopic scissors, and a bipolar coagulation electrode, we dissected the tumor and isolated the vascular pedicle, which we cauterized and resected. Then the neoplasm was completely removed via the foramen of Monro. Upon examination, the third ventricle was free of tumor tissue, and the ventricular aqueduct was well visualized with no evidence of occlusion (Fig. 2). Estimated blood loss was 15 ml. Analysis of the CSF collected intraoperatively revealed cytosis (1 neutrophil present), protein 11 mg/dl, chloride 115 mmol/L, and glucose 48.6 mg/dl.

Postoperative Course

The patient’s postoperative recovery was satisfactory. Brain MRI 3 days after the operation confirmed total resection of the tumor and a third ventricle size of 1.67 cm. There was still evidence of moderate periventricular edema around the anterior horns of the lateral ventricles (Fig. 3). At the time of discharge on postoperative Day 9, the patient was more active and without strabismus. Muscle strength was still +4 in the upper and lower extremities bilaterally with decreased muscle tone in all extremities. He had normal reflexes in the upper and lower extremities, with age-appropriate normal plantar reflex responses bilaterally. His appetite improved, and his head circumference had decreased by 1.5 cm. At the 10-month follow-up visit, there was no progression of the hydrocephalus and no evidence of tumor recurrence.

Pathological Findings

Histological examination revealed papillary structures with a large number of dilated blood vessels with hyperemia. The surface was lined with a single layer of cuboidal epithelium, which was morphologically consistent with a WHO Grade I choroid papilloma (Fig. 4).

Discussion

Meng et al. reported on the endoscopic removal of a cystic CPP of the third ventricle in an 18-year-old patient, with no evidence of recurrence during 2 years of follow-up observation.16 Reddy et al. described a combined approach...
endoscopic removal of a CPP in the third ventricle

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In half of all pediatric or adult cases, tumor is located in the lateral ventricle of the brain.

Eighty percent of third ventricular papillomas are located in the anterior region of the third ventricle, and a posterior localization of CPPs in the third ventricle is very rare.

According to Galassi et al., 14% of choroid plexus tumors occur during the 1st year of life. However, Nakano et al. described a CPP in the posterior aspect of the third ventricle in a 42-year-old woman, and Tomasello et al. treated a 49-year-old man with the same CPP localization.

Choroid plexus papillomas located in the third ventricle have their own characteristics. As the third ventricle is small and complex in its anatomy, it presents certain difficulties for surgical tumor removal.

Brain imaging shows CPPs to be round formations with tuberous surfaces. On both MRI and CT scans, CPPs

present as shapes with well-defined, large lobular margins. Computed tomography scans reveal a hyperdense structure. On MRI, T1-weighted sequences usually show a tumor that is isointense with the brainstem and T2-weighted sequences show a moderately hyperintense structure that can blend with the CSF in the image. Because the choroid papilloma is always highly vascularized, the injection of contrast leads to a notable increase in visualization. Tumor calcifications occur in 14%–25% of cases but are rarely seen in children.2,12,13,15 Blood supply to CPPs of the posterior areas of the third ventricles comes from the medial and lateral posterior choroidal arteries and the collicular arteries. Venous drainage is achieved via the internal cerebral veins, basal vein of Rosenthal, choroid veins (from the system of the vein of Galen), and anteromedial occipital veins.26

Because of the benign nature of the neoplasm and its low recurrence rates, patient survival rates after CPP resection are high (90%, 81%, and 77% for 1-, 5-, and 10-year survival rates, respectively).27 The most common postoperative complications are subdural hygroma and the need for CSF drainage. In a report by Pencalet et al., ventriculoperitoneal shunting was required in 50% of patients.20

Surprisingly, few reports in the literature discuss endoscopic CPP removal. We believe that current endoscopy equipment is technically adequate for handling such surgeries. Our experience with successful endoscopic removal of CPPs—not only from the third but also from one of the lateral ventricles—leads us to certain conclusions about the use of the neuroendoscopic approach. First, preoperative planning requires determining whether there is enough space around the neoplasm to allow safe surgical manipulations. To establish that, one must compare the relative volumes of the tumor and the ventricle as well as the size and type of endoscope (rigid, semirigid, flexible, channeled, and so forth). One must also assess the accessibility of the vascular pedicle by the endoscopic instruments and verify that it can be safely manipulated without causing damage to it or other neurovascular structures. Additionally, one must decide on the number of necessary access ports, for example, when the vascular pedicle is difficult to reach.

Moreover, the following factors are important to consider before the operation: 1) whether a bimanual technique must be used (that is, whether 2 instruments must be manipulated at the same time during a multiportal approach with multiple instruments); 2) whether it will be possible to change the trajectory and the angle of attack using different portals of access, if necessary; and 3) whether one can reliably clip and/or cauterize the vascular pedicle during the operation. Note that when dealing with a large tumor, we morcellate it and then remove it together with the port through the endoscopic hole of the working port.

**Conclusions**

Choroid plexus papillomas of the third ventricle are very rare compared with those in other locations. For this reason, even a single successful surgical treatment using a minimally invasive technique in a young pediatric patient is of great practical interest. Additionally, the clinical case described here demonstrates that a minimally invasive technique can be successfully used for complete monoporal endoscopic resection of a CPP in the posterior third ventricular regions.

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**FIG. 3.** Brain MRI with intravenous contrast after surgery. A: Sagittal section showing the tumor completely removed. Location of tumor prior to its removal (a). B: Sagittal section endoscopic hole (b) after transcortical monoportal access. Arrows indicate air in the anterior parts of the lateral ventricles. C: Coronal section of third ventricle (c), free of tumor. D: Axial section showing a decreased size of the third ventricle after tumor removal and regression of the internal hydrocephalus. Arrows indicate residual signs of the periventricular edema. Lumen of the third ventricle (d).

**FIG. 4.** Photomicrograph of a WHO Grade I CPP. H & E, original magnification ×10.
References

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Author Contributions

Conception and design: Gaibov, RA Sufianov. Acquisition of data: RA Sufianov. Analysis and interpretation of data: Gaibov. Drafting the article: Gaibov, RA Sufianov. Critically revising the article: all authors. Reviewed submitted version of manuscript: AA Sufianov. Approved the final version of the manuscript on behalf of all authors: AA Sufianov. Administrative/technical/material support: AA Sufianov. Study supervision: AA Sufianov.

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