Choroid Plexus Papillomas in two Siblings: Case Report

İki Kardeşte Görülen Koroid Pleksus Papillomu: Olgu Sunumu

ABSTRACT
Choroid plexus papilloma (CPP) is a rare, benign epithelial brain tumor of the nervous system seen particularly in infants. Familial cases are extremely uncommon. Some other form of malignant tumors was noted in the relatives of patients with CPPs, and some genetic defects regarding this coincidence were reported in the literature. These neoplasms are occasionally bilateral and hydrocephalus is an associated sign in most of the cases. We report three lateral ventricle CPPs in two siblings, at the age of 7 month and 2 years respectively. All tumors were resected with parietotemporal craniotomy and a superior temporal sulcus approach to the lateral ventricle. To avoid a concomitant need of ventriculoperitoneal shunt insertion, external ventricular drainage was inserted for a week in the postoperative period relieving symptoms of hydrocephalus. Search for a hereditary defect in the p53 gene of the second infant (7 months old) revealed no mutation. Postoperative courses were uneventful and the patients were followed for three years without any recurrence. Bilateral CPPs are rare and unusual in two siblings. A genetic predisposition such as the p53 mutation should be investigated in bilateral CPPs in particular.

KEY WORDS: Choroid plexus, Papilloma, Familiar, p53-Mutation, Lateral ventricle, Hydrocephalus

ÖZ

ANAHTAR SOZCÜKLER: Koroid pleksus, Papillom, Ailesel, p53 mutasyonu, Lateral ventrikül, Hidrosefali
INTRODUCTION

Choroid plexus papillomas (CPPs) are congenital neoplasms that are more frequently recorded in the pediatric population. They constitute 3% of all primary pediatric cerebral neoplasms (8, 22). Patients usually present with epileptic seizures and symptoms of hydrocephalus. They make up the most common ventricular tumor of childhood. About two thirds of choroid plexus papillomas are localized in the lateral ventricles (8,13,22). Genetic transition (3,11,17,19,23,24) or bilateral presentation is extremely rare and less than 10 cases have been reported previously (1,4-6,18). Here we report two siblings operated for unilateral and bilateral CPPs.

Case 1

A 2-year-old male infant was admitted with vomiting and epileptic seizures noted within the last 2 weeks. Neurological examination, clinical and family history were completely normal. An intensely enhancing mass in the right lateral ventricle trigone and hydrocephalus were noted on cranial CT. MRI showed a right lateral ventricle mass extending into the third ventricle and the preliminary diagnosis was CPP (Figure 1A). A right parietotemporal craniotomy was performed and the lesion was approached through the posterior portion of the superior temporal sulcus. A grayish-pink lobulated mass was removed grossly as total. An external ventricular drainage system was inserted in case of postoperative hydrocephalus. Drainage was removed a week later and the patient was discharged without any neurological deficit. Histopathological sections revealed typical findings of CPP. The antiepileptic drug was discontinued after 3-year seizure-free follow-up. MRI performed three years after surgery demonstrated total removal without any remnant and hydrocephalus (Figure 1B).

Case 2

A 7-month-old male infant experienced a seizure characterized with loss of consciousness and staring at a fixed point. His parents brought the child to the hospital since their elder son had been operated for the same symptom (Case 1). Further family investigation revealed no history of malignancy or a similar pathology. Neurological examination was unremarkable. CT demonstrated intensely enhancing bilateral lateral ventricle mass. On MRI, the lesions filled the lateral ventricles completely and hydrocephalus was prominent (Figure 2A). The infant was operated through superior temporal sulcus approaches bilaterally with an interval of 3 weeks between the operations. The excised lesions were put into liquid nitrogen directly after removal in order to perform genetic studies. Histopathological examination was again compatible with CPP. The patient was discharged 5 weeks after his admission without any neurological deficit. Imaging studies performed 2 years after surgery demonstrate complete removal of both tumors (Figure 2B). Study for a hereditary p53 gene mutation was negative shown by genetic analysis (Figure 3).

DISCUSSION

Choroid plexus papillomas are benign neoplasms which originate from neuroectodermal tissues. They constitute about 0.5 % of whole intracranial tumors.
Genetic transition and p53 mutation (location is at locus p53, 191170.0036) have infrequently been reported (17,23). Moreover, hSFN5/IN/1 somatic mutations of chromosome 22 were reported in CPP, atypical CPP and choroid plexus carcinomas were reported in the same family recently (19). CPPs in siblings or identical twins were reported twice previously (3,11). Other malignant tumors were noted in the relatives of patients with CPPs and some genetic defects regarding this coincidence were reported (6,19). We analyzed the p53 gene, which is a well known tumor suppressor gene, for the presence of germline mutations that can be associated with CPP. All coding exons of p53 were sequenced in both patients, which did not reveal any pathogenic mutations. An example of the sequence data where a previously reported mutation at codon 248 (CGG>TGG; Arg>Trp) resides is shown in Figure 3. Other tumor suppressor genes like hSFN5/IN/1 or oncogenes may have played a role in the development of the CPP in these two siblings.

Choroid plexus papillomas are usually localized in the lateral ventricles in children and fourth ventricle in adults (3,4,8,22). However, true bilateral lateral ventricle CPPs is rare (4,5,13,18). True bilateral CPPs in two siblings had not been reported until now. At the operation, we did not find any difference with regard to their surgical removal when compared to unilateral CPPs but one should be aware of this finding for operative planning. Another important point with regard to bilateral localization is to differentiate the contralateral papilloma from villous hypertrophy. Histopathological studies may be insufficient to differentiate these two entities and neuroradiological features may be used as an effective tool (7). Association of hydrocephalus, lobulated appearance and homogeneous enhancement are highly suspicious for CPPs.

Hydrocephalus is usually an associated feature of CPPs. In fact, the majority of tumors present with symptoms of hydrocephalus (6,14). Hydrocephalus in these patients is thought to be caused by an overproduction of CSF as noted in diffuse villous hyperplasia, and should thus resolve if tumor resection is complete (2). The role of inflammation and obstructive causes remains to be proved. Meticulous microsurgical techniques aid in total removal of tumor as well as prevention of hydrocephalus. The use of temporary external ventricular drainage also assists in prevention of postoperative hydrocephalus as indicated in this report. Drainage of blood and postoperative collections could also contribute to prevention of complications like hydrocephalus and subdural effusions (15). Shunting is an alternative strategy but should be avoided if possible since drop metastases of CPP are a well-known feature (9,10).

Surgical resection is the preferred way of treatment for CPPs. Several surgical approaches were defined previously for lateral ventricle CPPs (12,15,16,20). We prefer a transsulcal approach from the most posterior portion of the superior temporal sulcus. This approach provides a direct access to the posterior one-third of the lateral ventricle, called the trigone. The safety of the approach depends upon the cerebral tissue dissected although this region is anatomically safe, particularly on the non-dominant hemisphere. Tacconi et al. reported a good outcome if the resection was complete with a recurrence rate of less than 10 % (22). With regard to bilateral CPPs, we advocate a time interval of 3 weeks between the two operations if possible. There seems to be a chance of cure and low risk of recurrence for CPPs if the resection is complete. Role of radiotherapy in CPP treatment is still controversial and it is better to reserve this alternative just for recurrent and residual lesions with a risk of high morbidity and mortality (21). Follow-up is therefore the best choice to handle these patients.

REFERENCES


