Hemorrhagic Epidermoid Cyst in Cerebellar Vermis

Serebellar Vermis Yerleşimli Hemorajik Epidermoid Kist

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ABSTRACT

Intracranial epidermoid cysts are usually located in the paramedian regions with characteristic imaging features. Intracystic hemorrhage is rarely reported with most in the cerebellopontine angle area. We described a case of hemorrhagic epidermoid cyst in cerebellar vermis. The patient was a 21-year-old male presenting with a first episode of convulsive seizure attack as the initial and sole symptom. Head computed tomography showed a mass lesion in the cerebellar vermis with high density and nodular low density in the back of the lesion. Magnetic resonance imaging revealed most of the lesion was hypointense to isointense, extremely hypointensity on T1-, T2-weighted imaging respectively. The nodule was hyperintense on both T1- and T2-weighted images. The atypical clinical presentation, location and radiological features of intracranial epidermoid cyst make accurate diagnosis quite challenging. In such cases, scrutinized histopathological examination is necessary to exclude the malignancy that may need radio/chemotherapy.

KEYWORDS: Cerebellar tumor, Cerebellopontine angle, Epidermoid cyst, Hemorrhage

ÖZ


ANAHTAR SÖZCÜKLER: Serebellar tümör, Serebellopontin açı, Epidermoid kist, Hemoraji

INTRODUCTION

Intracranial epidermoid cyst represents approximately 0.2 to 1% of all intracranial tumors (3). They usually occur laterally. The most common location in the posterior fossa is the cerebellopontine angle (CPA) area. The fourth ventricle and cerebellar hemisphere location are also occasionally reported. However, occurrence in the cerebellar vermis is exceptionally rare (3, 4). We present a vermian epidermoid tumor with intracystic hemorrhage. The initial symptom was seizure and the radiologic imaging was unusual. We discussed the possible reasons for the atypical manifestations and emphasized the importance of histological examination.

CASE REPORT

A 21-year-old male soldier experienced a first episode of convulsive seizure attack when singing. On admission, he was alert without significant headache or cerebellar signs. Computed tomography (CT) of the brain disclosed a hyperdense mass lesion in the cerebellar vermis and obstructive hydrocephalus. A small portion within the lesion was found low in density (Figure 1A). Most of the lesion on magnetic resonance imaging (MRI) was hypointense to isointense on T1-weighted images and extremely hypointense on T2-weighted images (Figures 1B, C). Enhancement was not noticed (Figure 1D). The nodule in the back of this lesion was heterogeneously hypointense on both T1 and T2 weighted images, also absent of contrast enhancement.

He underwent midline suboccipital craniotomy. After opening the dura, an extra-axial mass with a transparent capsule was seen. We identified mud-like chronic hematoma in brown color and yellow lipid debris when the capsule was cut open (Figure 2A). During surgery, the lesion was totally removed; no hair or obvious calcification or ossification was observed. Postoperative course was uneventful. Histological examination showed keratin, stratified squamous epithelium and blood. Exhaustive looking for skin adnexal structures or tissue derived from mesoderm or endoderm as well as malignant elements failed. Epidermoid cyst accompanying hemorrhage was confirmed (Figure 2B). The patient recovered well with no morbidity. At 1-year follow up, MRI of brain revealed disappearance of hydrocephalus and the lesion.
DISCUSSION

Intracranial epidermoid cyst is a congenital lesion with a slow, long clinical course and an average age at diagnosis around 40 years (1). The CPA region is the most common location followed by the forth ventricle in the posterior fossa, where trigeminal neuralgia, hearing loss and gate ataxia are the main symptoms (1). The lesion in cerebellar vermis is extremely rare (1, 2). In our case, chronic hemorrhage may be responsible for the relatively young age at the disease onset.

Figure 1: A) Head CT showed a mass lesion in the cerebellar vermis with high density and nodular low density in the back of the lesion. B) On axial T1-weighted MRI, the 53 ×55×48mm lesion was hypointense to isointense and the nodule was hyperintense. C) Axial T2-weighted MRI shows extremely hypointensity of the lesion and the nodule is heterogeneously hyperintense. D) No enhancement is observed on sagittal contrast T1-weighted MRI.

Figure 2: A) Photograph during operation revealed mud-like old hemorrhage and yellow lipid debris (arrow). B) Histological examination showed keratin, stratified squamous epithelium and blood, consistent with hemorrhagic epidermoid cyst (HE, ×200).
onset due to accelerated growth of the lesion. In addition, seizure attack became the single presentation, probably induced by singing which contributed to the acute elevation of intracranial pressure in the setting of hydrocephalus. It was different from the report by Singh et al. in which the patient presented seizure in sleep because of acute hemorrhage of epidermoid cyst in the supratentorial compartment (5).

The radiological manifestation of our case was quite atypical. It has been reported that intratumoral hemorrhage is the most common cause of changes in the signal intensity (1, 4). Meanwhile, the signal intensity on MR imaging varies depending on the contents. Hyperintensity on both T1- and T2-weighted images can be due to a semi-liquid cystic content with high protein concentration; high signal intensity on T1-weighted and low signal intensity on T2-weighted images can be caused by even thicker toffee-like content with a combination of high protein content and high viscosity; low signal intensity on T1- and T2-weighted images can be attributed to an unusual composition of epithelial debris, crystals of cholesterol, and keratin (1). In our case, the signal intensity of most of the lesion partly accords with that of chronic hematoma while the signal feature of the nodule may be related to the intracystic content of high protein.

When epidermoid cyst occurs in the cerebellar vermis with hemorrhage, preoperative misdiagnosis is quite frequent (1, 4). Surgeons should recognize its uncommon features to avoid surgical complications such as chemical meningitis. Additionally, careful histological examination is essential to differentiate from other diseases such as immature teratoma and to detect malignant transformation which requires adjuvant therapy after surgery (2).

CONCLUSION

Vermian epidermoid cyst is very rare and can have seizure as the sole clinical presentation. When complicating with hemorrhage, the differentiation from other entities is difficult by preoperative imaging features. Scrutinized histopathological examination is necessary to exclude the malignancy which may need radio/chemotherapy.

REFERENCES