A Patient with a Dysembryoplastic Neuroepithelial Tumor Who Underwent Epilepsy Surgery after Initial Seizure

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Key Words
Dysembryoplastic neuroepithelial tumor · Cortical dysplasia · Epilepsy surgery · Electrocorticography · Initial seizure

Abstract

Background: Since dysembryoplastic neuroepithelial tumors (DNTs) are benign tumors that are frequently associated with long-standing medically intractable epilepsy, it is well known that the surgical strategy is resection of the associated epileptogenic zone as well as the tumor. However, the surgical strategy for DNT with a single seizure has not been fully discussed. Methods: We report an 8-year-old boy with DNT in the nondominant frontal lobe who underwent epilepsy surgery at 3 months after his initial seizure. Results: An intraoperative electrocorticogram revealed frequent paroxysmal cortical activity lateral to the tumor. Since resection of the tumor resulted in persistent paroxysmal activity in this cortex, additional resection was performed. The histological findings in the cortex revealed the presence of cortical dysplasia (CD) (Palmini type IIA). Lesionectomy alone might have left the epileptogenic CD. Conclusion: It is thought that epilepsy surgery should be recommended in patients with typical neuroimaging findings of DNT, even if the patients had only one episode of seizure.

Introduction

Dysembryoplastic neuroepithelial tumors (DNTs) are glioneuronal tumors that are frequently associated with ‘long-standing’ medically intractable focal epilepsy in children and young adults, and were initially described by Daumas-Duport et al. [1]. Since DNTs are benign tumors, the aim of surgery is to treat the associated intractable epilepsy, and it is well known that the surgical strategy is resection of the associated epileptogenic zone and the tumor, which are identified by chronic and/or intraoperative electrocorticography (ECoG) [1–16]. However, the surgical strategy in DNT with a single seizure or well-controlled seizures has not been fully discussed. We report a patient with DNT who underwent epilepsy surgery at 3 months after the initial seizure.
Case Report

An 8-year-old right-handed boy suddenly squatted while dancing as part of his school work and subsequently developed a generalized tonic seizure. He was transferred to the pediatric department of the local hospital. An electroencephalogram (EEG) demonstrated frequent paroxysmal discharges in the right frontal region which were treated with 400 mg of valproate. Since a computed tomography (CT) scan revealed a tumor in the right frontal lobe, he was referred to us for tumor resection.

Upon admission, the neurological findings were normal. Although his seizure was completely controlled by valproate monotherapy, the EEGs still revealed frequent paroxysmal activity in the right frontal region (F4 of the 10–20 International System). A CT scan demonstrated a low-density tumor in the right frontal lobe (fig. 1a). Deformity of the skull overlying the tumor is noted. No gadolinium enhancement was noted (fig. 1b, c).

At 3 months after the initial seizure, a right frontal craniotomy was performed to confirm the histological diagnosis of the tumor. In addition to tumor surgery, epilepsy surgery monitored by intraoperative ECoG was indicated since reduction or cessation of postoperative antiepileptic drug treatment was desired by his parents. Although the tumor was not exposed on the cortex, its location was identified by surface anatomy MR scanning (fig. 2a, b) [17, 18]. The gyrus above the tumor was enlarged. An intraoperative ECoG was recorded from the cortex and demonstrated frequent paroxysmal activity, especially over the cortex, just lateral to the tumor (ECoG irritative zone) (fig. 2c). There was a sulcus between the tumor and the ECoG irritative zone (fig. 2b). Resection of the gyrus involving the tumor was performed. Since the ECoG recorded from the cortex around the tumor cavity showed that there was still frequent paroxysmal activity on the ECoG irritative zone lateral to the tumor location, additional resection of this cortex was performed (fig. 2d). The final ECoG demonstrated disappearance of the paroxysmal activity in the right frontal lobe (fig. 2e).

Fig. 1. a Serial axial images of a CT scan demonstrating a low-density tumor in the right frontal lobe. Deformity of the skull overlying the tumor is noted. b, c Coronal (b) and sagittal (c) views of T1-weighted images following gadolinium administration depict a low-intensity tumor in the cortex of the right frontal lobe. A pseudocystic appearance is observed. No gadolinium enhancement is noted. d Serial axial views of T2-weighted images demonstrate a high-intensity tumor in the right frontal lobe.
Fig. 2. a The right anterior quadrant of a surface anatomy MR image clearly demonstrates the anatomical relationship between the low-intensity tumor and the surrounding gyri, sulci and cortical veins. The numbers of the recording electrodes are indicated on this image to show the spatial relationship between the tumor and the recording electrodes. b An intraoperative photograph, the orientation of which is matched with a, demonstrates that the gyrus above the tumor (T) is enlarged, although the tumor is not exposed on the cortex. The white arrows indicate the sulcus between the tumor location and the irritative zone on the intraoperative ECoG (asterisks). c Intraoperative ECoG depicting frequent paroxysmal discharges not on the tumor but on the gyrus just lateral to the tumor (electrode No. 13 and 14). d Intraoperative photograph taken after removal of the tumor showing the special relationship between the surgical defect (white arrows) and the ECoG irritative zone (asterisks). e The final ECoG demonstrates disappearance of the paroxysmal discharges in the right frontal lobe.
The postoperative course was uneventful. With a tapering dose of valproate, the boy was seizure free during the initial 2 years postoperatively. A postoperative EEG showed disappearance of the paroxysmal activity in the right frontal region.

The histopathological findings (hematoxylin and eosin, HE, staining) of the tumor were consistent with those of DNT. The tumor exhibited a multinodular architecture leading to expansion of the cortex. The nodular focus of the tumor was well delineated from the cortex (fig. 3a). The tumor displayed specific glioneuronal elements, such as oligodendroglia-like cells and floating neurons in a myxoid matrix (fig. 3b). The additionally resected ECoG irritative zone exhibited dyslamination of the neurons (fig. 3c) and dysmorphic neurons with clumped Nissl bodies (fig. 3d). Since no balloon cells were present, the findings indicated cortical dysplasia (CD) of Palmini type IIA [19].

**Discussion**

Histopathologically, DNT is a benign, predominantly intracortical lesion mainly composed of a population of oligodendrocyte-like cells with admixtures of mature ganglion cells and astrocytes that are located in a myxoid or dense neurofibrillary matrix [1, 5]. Involvement of the cortical gray matter and a multinodular growth pattern are typical findings [1, 5, 20, 21]. On the basis of its histopathological appearance and benign clinical course, it has been demonstrated that the typical radiological features of DNT include a cortical hypodense lesion with deformity of the calvaria adjacent to a superficially located tumor on CT scans and a T₁- and T₂-prolonged tumor with a multicystic appearance on MRI [1, 5, 13, 20–23].
sions characteristically lack surrounding edema and only a few are enhanced after addition of contrast material [1, 5, 20, 21, 23]. The neuroimaging findings in the present case were consistent with those of DNT. However, these reported neuroimaging features are nonspecific and may be seen in a variety of low-grade tumors, particularly gangliogliomas, gangliocytomas, and low-grade astrocytomas [13]. Although functional neuroimages, such as fluorodeoxyglucose and methionine positron emission tomographies (PET) and proton magnetic resonance spectroscopy, especially high methionine uptake on PET, could contribute to do a differential diagnosis from low-grade astrocytomas, these findings are not conclusive [22, 24]. Thus, in the present case, tumor surgery was indicated to confirm the histological diagnosis of the tumor.

There is no consensus thus far regarding the timing of the surgery. In a series described by Ostertun et al. [21], one of 16 epileptic patients with DNT underwent surgery at 0.2 years after the onset of epilepsy. Fernandez et al. [20] reported 14 DNT patients, and described that the preoperative duration of epilepsy in 7 patients was 4–9 months. In a recent report by Minkin et al. [10], 12 of 24 DNT patients did not have pharmacoresistant epilepsy. Furthermore, the preoperative minimal duration of epilepsy was reported to be 0.25 years. Thus, it seems that surgical resection of DNT is indicated, regardless of whether or not the patient has ‘long-standing’ intractable seizures.

With regard to the surgical strategy for control of the ‘long-standing’ intractable epilepsy associated with DNT, a few authors previously demonstrated that lesionectomy alone of a DNT lesion achieved good outcomes for seizure control [6, 10]. However, recent studies clearly showed that DNT epileptogenicity is not intrinsic but instead is located in a histologically dysplastic area of the cortex, since DNT is frequently associated with CD [1, 5, 14, 16, 22–27], which has intrinsic epileptogenicity [2, 17]. Recently, Lee et al. [8] demonstrated that amplification of ionotropic glutamate receptors such as GluR2 and GluR3 in the associated CD lesion might be the underlying cause of epileptic seizures. Therefore, it seems necessary to resect the DNT with the associated CD for effective seizure control [3, 7, 9, 14, 16].

However, associated small CD lesions are still quite difficult to detect by preoperative neuroimaging examinations, even though the currently available high-resolution MRI techniques have improved the detection of CD compared with past decades [2, 9]. Therefore, a physiological method that can localize an epileptogenic CD lesion is considered to be important [4, 9, 15, 22, 23]. Although many previous authors reported that the CD in DNT patients is focally adjacent to the tumor [1, 4–6, 14, 15, 22–27], there are surprisingly few reports regarding the topographical relationship between the epileptogenic cortex and the CD location [8, 9, 16]. Mikuni et al. [28] clearly demonstrated a relationship between the ECoG irritative zone and the type Ib (not type II) CD location in a patient with DNT.

In epilepsy surgery, invasive video-intracranial EEG monitoring with subdural grid electrodes is still a gold standard to aid in the localization of the epileptogenic zone and the functionally eloquent zone [17, 18]. However, in the present case, the tumor location was the nondominant frontal lobe, and placement of a chronic intracranial electrode was too invasive for an 8-year-old boy with only one episode of epilepsy. Thus, epilepsy surgery under the guidance of an intraoperative ECoG was selected according to many previous reports [4, 8, 9, 11, 15, 22, 23, 28]. Although there is no consensus regarding the indication for epilepsy surgery in patients with only one episode of seizure, epilepsy surgery with an intraoperative ECoG could be performed in the present case without additional invasiveness during the tumor surgery.

In our case, the intraoperative ECoG clearly depicted the epileptogenic cortex lateral to the tumor and the histological examination showed a CD lesion in the ECoG irritative zone. There was a sulcus between the tumor and the epileptogenic CD lesion and no continuity was noted between the tumor and the CD lesion. Lesionectomy of the tumor (tumor surgery) alone left behind an epileptogenic CD in another cortex, which might serve as a focus of recurrent or continued epilepsy [4, 8, 9, 15, 22, 23, 27, 28]. Although a long-term follow-up should be performed in the present case, he was seizure free during the first 12 months postoperatively with a tapering dose of antiepileptic drug, and antiepileptic drug cessation is hoped for in the near future. Although a firm conclusion cannot be drawn from a single case report, epilepsy surgery is recommended in patients with typical neuroimaging findings of DNT, even if they have only one episode of seizure.

References


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