SURGICAL AND SEIZURE TREATMENT OUTCOMES IN ADULT DYSEMBRYOPLASTIC NEUROEPITHELIAL TUMORS: A CASE SERIES

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SUMMARY – Dysembryoplastic neuroepithelial tumors (DNETs) are benign neoplasms classified in the category of glioneuronal tumors. The estimated incidence of DNETs is 0.03 per 100,000 person per year with the age peak in a range between 10 and 14 years, and decreasing dramatically with increasing age. They are seldom diagnosed in persons above 20 years of age, being a cause of tumor-related intractable epilepsy that begins in childhood or adolescence. They have been proven to be the second most common type of epileptogenic tumors in pediatric population. These rare tumors cause chronic drug-resistant partial complex seizures with or without secondary generalization. Herein, we provide institutional case series of six adult patients with temporal lobe DNET presenting with complex partial seizures. Lesionectomy was performed with tumor resection in toto in three patients. In another three, partial resection was performed, whereas tumor remnant was left intact to avoid possible basal ganglia damage. All patients were seizure free postoperatively. Lesionectomy alone in temporal lobe epilepsy was associated with less favorable outcome than anterior temporal lobectomy. Total tumor removal is considered a major prognostic factor in most studies.

Key words: Dysembryoplastic neuroepithelial tumor; Seizure; Surgery; Outcome

Introduction

Dysembryoplastic neuroepithelial tumors (DNETs) are benign tumors first described by Daumas-Duport et al.1. The World Health Organization (WHO) includes DNETs in the category of neuronal and mixed neuronal–glial tumors, corresponding to WHO grade 12. Four DNET histologic subtypes have been described3-6. They represent 1.2% of all neuroepithelial tumors resected in patients under the age of 20 years7. In epilepsy surgery series, the prevalence of DNETs ranges from 0.8% to 19%8-10. DNETs are found in about 20% of histologic diagnoses in adult epilepsy surgery centers11. The most common seizure type with DNETs is complex partial seizures with epileptogenic zone in the temporal lobe12. Surgery is associated with very good prognosis and low recurrence risk8. Lesionectomy alone in temporal lobe epilepsy is associated with less favorable outcome than anterior temporal lobectomy.

Herein, we describe institutional series of six adult patients with such tumors presenting with complex partial seizures, and discuss postoperative surgical and seizure outcome.
Six adult patients with DNETs admitted to Department of Neurosurgery, Sestre milosrdnice University Hospital Center during the 2012-2018 period were included. Patient characteristics are summarized in Table 1. All patients were right-handed and their mean age was 29.8 (range, 19 to 43) years. All patients underwent presurgical epileptogenic, neuroradiological and neurophysiological examination. All patients had complex partial seizures with a mean duration of 13.2 years. Four patients were on one antiepileptic drug (AED) and two patients on two AEDs prior to surgery. Preoperative neurophysiological examination revealed normal results in four younger patients, but reduced verbal memory in two older patients.

Four patients had DNET in the left temporal lobe and two in the right temporal lobe. In all patients, DNET involved the uncus and amygdala without hippocampus involvement (Fig. 1A, B).

Surgery was performed and tumor resection in toto was achieved in three patients (Fig. 2). In another three patients, subtotal resection was performed in order to avoid basal ganglia damage. Postoperative period was uneventful, without complications.

Histopathologic examination showed complex DNET type in all six cases. Follow-up was 24 months.

Table 1. Patient characteristics

| Case | Sex | Age (yrs) | Location of DNET | Seizure type | No. of AEDs | Seizure | Surgery | Preop/postop outcome |
|------|-----|-----------|------------------|--------------|------------|---------|---------|----------------------|
| 1    | M   | 20        | Left temporal    | CP           | 1/0        | SF      | Lesionectomy |                      |
| 2    | M   | 27        | Left temporal    | CP           | 1/1        | SF      | Lesionectomy |                      |
| 3    | F   | 30        | Left temporal    | CP           | 2/1        | SF      | Lesionectomy |                      |
| 4    | F   | 19        | Right temporal   | CP           | 1/0        | SF      | Lesionectomy |                      |
| 5    | M   | 43        | Left temporal    | CP           | 2/2        | SF      | Lesionectomy |                      |
| 6    | M   | 40        | Right temporal   | CP           | 2/2        | SF      | Lesionectomy |                      |

DNET = dysembryoplastic neuroepithelial tumor; AEDs = antiepileptic drugs; CP = complex partial; SF = seizure free
Postoperatively, all six patients were seizure-free, and AEDs were withdrawn in two youngest patients.

Discussion

Epilepsy onset in late childhood, complex partial seizures concordant with tumor location, and male predominance are characteristic of DNET\textsuperscript{13,14}. The typical magnetic resonance imaging (MRI) pattern consists of a pseudocystic or multicystic appearance, strongly hypointense on T1-weighted and hyperintense on T2-weighted images\textsuperscript{15}. Tumor subtypes may be recognizable on MRI allowing simple and complex histologic forms to be differentiated from nonspecific forms\textsuperscript{14}. In our case series, all six patients were type 1 according to MRI classification.

Surgical series report favorable outcomes in 70% to 90% of cases\textsuperscript{1,3,8,10-17}. Surgical methods consist of lesionectomy or corticectomy, including amygdalo-hippocampectomy and anterior temporal lobectomy. In our six-case series, we performed lesionectomy, including removal of the uncus and temporal ventricular part of amygdala that were involved by the tumor. We noticed that all lesions were sharply delineated from the hippocampus and could be dissected leaving the hippocampus intact. We found that important for patient postoperative neurophysiological functioning, although we did not perform postoperative neurophysiological examination.

Whether surgical outcome is related to the size of resection or type of the procedure remains controversial. Complete tumor removal is considered a major prognostic factor in most studies\textsuperscript{10,12,14,15,17}. Incomplete resection is identified as the main cause of surgical failure\textsuperscript{12}. In our three patients, we did not resect the part of the lesion in the extra-ventricular portion of amygdala towards the basal ganglia in order not to damage basal ganglia. Nevertheless, all three patients were seizure-free postoperatively.

Surgical failure has also been attributed to the presence of dysplastic cortex adjacent to the tumor. Removing these areas has been considered necessary for a favorable outcome\textsuperscript{8,12}, but this has been contradicted by others\textsuperscript{10,16}. Lesionectomy alone in temporal lobe epilepsy has been associated with a less favorable outcome than anterior temporal lobectomy\textsuperscript{17}. This may be explained by the network organization of the epileptogenic zone and hippocampal involvement in tumors with temporal location\textsuperscript{5,14}. Although we performed lesionectomy in all six patients, seizure outcome was favorable in all cases.

The second major prognostic factor has been related to young age at surgery and short epilepsy duration\textsuperscript{10,14,16,17}. In our series, median epilepsy duration was 13.2 years due to the fact that we had two adults aged 43 and 40 years with long epilepsy duration. In contrast, another four patients were younger and were seizure-free even after AED withdrawal. This is in accordance with the finding reported by Fay-McClymont et al. that successful AED discontinuation was also related to early surgery in young subjects\textsuperscript{18}.

No tumor recurrence was noticed during the follow-up in our series, despite those three patients with tumor remnants. In their large series, Campos et al. found that recurrence was rare and occurred at the site of residual tumor\textsuperscript{15}. These findings advocate performing complete tumor resection whenever possible.

In conclusion, DNETs are found in about 20% of the histologic diagnoses in adult epilepsy surgery centers. Complete tumor removal is considered a major prognostic factor. In our case series, all patients underwent lesionectomy leaving the hippocampus and para-hippocampus intact with favorable seizure outcome achieved in all cases.

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ISHOD KIRURŠKOG LIJEČENJA I EPILEPTIČNIH NAPADAJA  
KOD DISEMBRIOPLASTIČNIH NEUROEPITELIJSKIH TUMORA U ODRASLIH:  
PRIKAZ SERIJE SLUČAJEVA

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Disembrioplastični neuroepitelijski tumori (DNET) dobroćudne su novotvorine koje se svrstavaju u skupinu glioneuralnih tumora. Njihova je godišnja učestalost 0,03 na 100.000 stanovnika s vrškom pojavnosti u dobi između 10 i 14 godina života. Učestalost se smanjuje s porastom dobi, a tumori se rijetko javljaju nakon 20. godine života. Dokazani su kao uzrok epilepsije koja započinje u djetinjstvu i adolescenciji te kao drugi najčešći tip tumora koji uzrokuje epileptične napadaje u dječjoj dobi. Ovi rijetki tumori uzrokuju kronične i na terapiju otporne parcijalne složene epileptične napadaje s generalizacijom ili bez nje. U ovom radu prikazujemo institucijsku seriju šest odraslih bolesnika s DNET-om u području temporalnog režnja koji su se prezentirali složenim parcijalnim epileptičnim napadajama. Tri su bolesnika kirurški liječena lezionatektomijom s totalnom resekcijom tumora. U preostalih tri bolesnika učinjena je parcijalna resekcija, a dio tumora je ostavljen kako bi se izbjeglo oštećenje bazalnih ganglija. Svi su bolesnici nakon zahvata bili bez epileptičnih napadaja. U slučaju temporalne epilepsije, a u usporedbi s prednjom temporalnom lobektomijom, selektivno uklanjanje tumorske lezije bilo je povezano s lošijim ishodom liječenja. Uklanjanje tumora u cijelosti smatra se važnim prog nostičkim pokazateljem ishoda liječenja u većini provedenih istraživanja.

Ključne riječi: Disembrioplastični neuroepitelijski tumori; Epileptični napadaji; Kirurško liječenje; Išod liječenja