Juvenile Myoclonic Epilepsy: Myoclonic Status Epilepticus without Coma - Report of Three Cases

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Status epilepticus (SE) is rare in juvenile myoclonic epilepsy (JME). This report presents three patients with myoclonic status epilepticus (MSE). MSE is defined as prolonged period of myoclonic jerks that are correlated with epileptiform discharges on electroencephalogram. The precipitating factors among the three patients were: introduction of carbamazepine in case1, missing the dose in case2, and introduction of oxcarbazepine in case3. Of the three patients, one patient was a misdiagnosed case of JME. In him the diagnosis of JME was established after 35 years when he developed MSE with the addition of oxcarbazepine to the antiseizure medication (ASM) which he was taking. Detailed review of the history revealed that he used to get occasional myoclonic jerks with deprived sleep and stress. This patient illustrates that the diagnosis of JME can be missed or delayed if history of myoclonic jerks is not elicited, particularly in patients with pubertal onset epilepsy. The other lesson is that possibility of JME should be considered in patients with drug resistant epilepsy (pseudo-drug resistance). (2020;10:92-95)

Key words: Juvenile myoclonic epilepsy, Misdiagnosis, Myoclonic status epilepticus, Oxcarbazepine

Introduction

Juvenile myoclonic epilepsy (JME) is a clinically heterogenous, generalized epilepsy syndrome with peri-pubertal onset. Myoclonic seizure is the defining seizure type of the syndrome and other seizure types include generalized tonic-clonic seizures (GTCS) and absences. Myoclonic jerks and GTCS typically occur on awakening. Photosensitivity, eye closure sensitivity, orofacial reflex myoclonia, and praxis induction are the typical epileptic traits of JME. The diagnostic electroencephalogram (EEG) findings include normal background activity with generalized 3-6 Hz spike-waves with frontal predominance. Treatment response to valproate (VPA) is generally good. Status epilepticus (SE) is rarely reported in patients with JME. In a study of 133 patients with JME, only five (3.8%) were diagnosed with SE.3 The most frequently reported SE subtypes are myoclonic SE (MSE) and nonconvulsive SE (NCSE) or absence SE (ASE). Very rarely it can be mixed absence-myoclonic status. MSE is defined as prolonged period of myoclonic jerks that are correlated with epileptiform discharges on EEG. We report three patients with JME who developed MSE.

Case Report

In this study the diagnosis of JME was based on the criteria proposed by the international experts on JME.4 The clinical details of the patients are given in Table 1. The SE subtype was MSE in all the three patients and they were conscious throughout the period of SE. The time delay in the diagnosis of SE was mainly due to the time taken by the patients to arrive at emergency department from their place of domicile. Myoclonic jerks were brief, synchronous, grossly symmetric, and involved more upper limbs. In case1 the jerks were frequent and involved both distal proximal muscle of both the upper limbs making her prone to dropping things. Case1 was on an inappropriate antiseizure medication (ASM) and carbamazepine (CBZ) when presented with MSE. The female patient was diagnosed with JME, well controlled on VPA (600 mg per day). She stopped valproate (VPA) after her marriage and had an episode of GTCS for which she was prescribed CBZ by the family physician.

Case3 was a misdiagnosed case of JME. The diagnosis of JME was established only when the male patient developed MSE with the use of oxcarbazepine (OXC). He consulted first time the physician for the new-onset GTCS at the age of 20 years and was initially put on phenobarbital and phenytoin with which he used to have GTCSs once in 1-2 months. He consulted several physicians including neurologist and was extensively investigated with 3-T magnetic resonance imaging and multiple EEGs which were all normal. He was tried on several ASM. When he consulted us, he was on phenobarbitone and...
Table 1. Clinical characteristics of patients JME and myoclonic status epilepticus

| Age/sex | Primary diagnosis | Precipitating factor | Status type and clinical features | EEG features | Treatment and outcome |
|---------|-------------------|----------------------|-----------------------------------|--------------|-----------------------|
| Case 1: 25 years/female | Diagnosed case of JME and on valproate 600 mg, counselled regarding marriage and pregnancy | Stopped valproate after her marriage, had an episode of GTCS, local physician changed valproate to carbamazepine | MSE (duration 16-17 hours) Conscious, no neurological deficits, frequent myoclonic jerks predominantly distal involving proximal and distal muscles of upper limbs, occasional dropping of objects from hands | Ictal EEG: frequent paroxysmal generalised poly-spike and occasional poly-spike-wave discharges, almost continuous 4-5 Hz (Fig. 1A) | MSE was terminated with IV midazolam (0.2 mg/kg) followed by valproate 40 mg/kg loading dose and was put on valproate 600 mg per day. In the follow-up valproate was changed to levetiracetam 1,000 mg/day and was counselled regarding conception and pregnancy |
| Case 2: 20 years/male | JME on valproate 1,000 mg/day, no seizures | Stopped valproate since one month as he is seizure free | MSE (duration 16-18 hours) Conscious, no neurologic deficits Frequent myoclonic jerks mainly involving proximal muscle in both upper limbs | Ictal EEG: frequent paroxysmal generalized spike and wave and occasional poly-spike-wave discharges, 3-4 Hz (Fig. 1B) | MSE was terminated with IV midazolam (0.2 mg/kg) followed by valproate 40 mg/kg loading. He was continued on valproate 500 mg q12h |
| Case 3: 55 years/male | Misdiagnosis of JME and being treated as a case of drug resistant epilepsy with phenobarbitone and levetiracetam | He visited our clinic, we also considered the diagnostic possibility of drug resistant epilepsy and oxcarbazepine was added, following which he developed frequent myoclonic jerks | MSE (duration 48 hours) Conscious, no neurologic deficits, frequent jerking of both the upper limbs mostly proximal muscles | Ictal EEG: frequent paroxysmal generalized poly-spike and rare spike-wave discharges, 4-5 Hz (Fig. 1C) | MSE was terminated by IV midazolam 0.2 mg/kg followed by IV loading dose of valproate (40 mg/kg). He was started on valproate 1,000 mg q12h and the other ASMs are being withdrawn |

JME, juvenile myoclonic epilepsy; EEG, electroencephalogram; GTCS, generalized tonic-clonic seizures; MSE, myoclonic status epilepticus; ASM, antiseizure medication.
Figure 1. (A) Ictal EEG in case 1. Anterior-posterior longitudinal bipolar montage showing normal background activity and frequent paroxysms of generalized symmetrical and synchronous poly-spikes and occasional poly-spike and wave discharges. Patient was fully aware during the EEG recording and having myoclonic jerks. (B) Ictal EEG in case 2. Anterior posterior longitudinal bipolar montage showing normal background activity and frequent bursts of generalized, symmetrical, and synchronous 3-3.5 Hz spike-wave and poly-spike and wave discharges. Patient was aware during the EEG recording and having myoclonic jerks. (C) Ictal EEG in case 3. Anterior posterior longitudinal montage showing normal background activity and frequent bursts of generalized, symmetrical, and synchronous poly-spike and spike-wave discharges. Patient was aware during the EEG recording and having myoclonus. EEG, electroencephalogram.
annual incidence of NCSE were 5.8% and 1.2%, respectively. The new classification of SE categorizes the status into CSE and NCSE based on the presence or absence of prominent motor symptoms and the degree of impaired consciousness. MSE is categorized under convulsive SE and is further divided into MSE with coma (mixed absence myoclonic SE) and MSE without coma. NCSE or ASE is categorized under NCSE without coma, typical absence status. In the series reported by Larch et al., three patients had MSE immediately after termination of GTCS and two patients had GTCS with myoclonic absence. In our patients, MSE was the SE subtype.

The new definition of SE is conceptual, with two time points: \( t_1 \) (when a seizure is likely to be prolonged leading to continuous seizure activity) and \( t_2 \) (when a seizure may cause long-term consequences). The \( t_1 \) for ASE is considered to be 10-15 minutes and time for \( t_2 \) is uncertain. For MSE the data are not yet available to define the two time points. In view of these limitations, we define MSE as “continuous or intermittent bilateral myoclonic jerks with preserved awareness lasting for at least 30 minutes”. In case1 and case2, the duration of status was about 16-18 hours and in case3 it lasted for 48 hours. A comparative study for differences between JME with SE and JME without SE observed that myoclonic jerks limited to specific body parts (one arm, face or head) were significantly more common in patients with SE.2

The common and important precipitants for SE in patients with JME are deprived sleep, inappropriate ASM, ASM noncompliance, and alcohol.2-5 The precipitant in case1 was inappropriate ASM, in case2 ASM nonadherence, and in case3 misdiagnosis and use of inappropriate ASMs, OXC. The delay in diagnosis was due to failure to ask for myoclonic jerks, even he had new-onset GTCS around pubertal age. The correct diagnosis was made only after he developed MSE with the use of OXC. Aggravation of genetic generalized epilepsy syndromes by inappropriate ASMs has been well documented, particularly with CBZ. Precipitation of MSE and ASE by inappropriate use of CBZ in JME has rarely been reported.10 Use of OXC in JME can exacerbate myoclonic and absence seizures.11 The first report of SE in JME with the use of OXC was done by Fanella and colleagues. This report is probably the second patient. He is a misdiagnosed case of JME and precipitation of MSE with the use of OXC established the diagnosis of JME after 36 years of onset of epilepsy. Jeong et al.13 reported two patients diagnosed with JME by first-ever status epilepticus in adult life. These three patients’ cases emphasizes the importance of asking for myoclonic jerks in patients with new-onset GTCS around pubertal age. GTCS is the seizure type for which most patients with JME seek medical attention for the first time.

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