COMPLEX PARTIAL STATUS EPILEPTICUS IN A CHILD

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SUMMARY - Complex partial status epilepticus (SE) has been reported rarely in children. We describe the clinical case of a 14 year-old girl with complex partial seizures (CPS) since age 10 who developed a complex partial SE probably because she was not adherent to treatment. The neurologic examination and computed tomography scan were normal. During the SE she received diazepam and phenytoin and became free of the seizures after 5 minutes. The ictal EEG showed spikes and slow waves over the right temporal region.

KEY WORDS: complex partial status epilepticus, EEG, child.

CASE REPORT

ASS, a female child, was first seen at the age of 14 years when she was referred with a history of numerous daily attacks of blank spells. She was hospitalized for investigation and treatment. The antenatal and birth histories had been unremarkable. There was no history of consanguinity or family history of seizures. In retrospect, the parents felt that she had been having staring spells from the age of 10 years when the treatment begun but she was not adherent. These spells lasted less than 3 min and were associated with alimentary (chewing, lip and tongue smacking, swallowing, or licking one’s lips) and gestual automatisms. Psychomotor development was normal. Her performance in school was good. During the examination she used to stare with fixed gaze and frozen face, periods of confusion and we observed characteristic automatisms that consisted mainly of grasping objects and frightened expression.
Fig 1. EEG carried out during the complex partial status of our patient: continuous slow waves intermixed with spikes and sharp discharges over the right fronto-temporal region and with spread to the contralateral side.

Fig 2. Interictal EEG: isolated and independent sharp waves over both anterior temporal regions.
A conventional EEG (CEEG) was recorded for 30 min and showed continuous discharges characterized by slow waves intermixed with spike and sharp components in various degrees over the right fronto-temporal region with spread to contralateral region (Fig 1). A diagnosis of complex partial SE was made, and she was treated with diazepam and phenytoin. Clinical events were not observed after initiation of treatment. A second CEEG was recorded, at which time she had been seizure-free for 2 hours. This recording showed a normal background activity and interictal paroxysmal discharges by isolated sharp waves independently over both fronto-temporal regions but with a predominance on the left side (Fig 2). Physical and neurological examination were normal. Computed tomography scan was also normal.

Following discharge from hospital, the girl was seen at regular intervals as an outpatient and, at age 15, while on carbamazepine therapy, she improved of the seizures, exhibiting a frequency of 12 to 16 per year.

COMMENTS

The variable clinical expression and the need of EEG for confirmation become very difficult, sometimes, the recognition of complex partial SE. The first case was published by Gastaut and Roger (1956) and has previously been considered a rare phenomenon, but the actual incidence of this disorder may be underestimated. Temporal lobe SE and psychomotor SE are the other terms that have been used to describe this phenomenon, but is not an appropriate designation, since seizures emanating from extratemporal regions may be associated with complex partial SE.

Several criteria for diagnosis of complex partial SE have been proposed. Mayeux and Lueders stressed focal EEG epileptic discharges, either continuous or recurrent, while Treiman and Delgado-Escueta emphasized recurrent or cyclical behavioural seizures. Our patient illustrates well the continuous type. It is our opinion that this classification into discontinuous and continuous forms is purely descriptive, and transitions from one to the other type may exist.

The ictal EEG in our patient showed discharges arising in the right temporal region, confirming the diagnosis of complex partial SE. The majority of the previously reported cases were, also, of temporal origin, but a few had an occipital focus. Moreover the features of complex partial SE were consistent in our patient. The patient had impaired consciousness characterized by absence of vocalization, lack of interaction with familiar people, decreased reaction to pain, and intermittent picking at clothing or nearby objects. The episode was terminated after 5 minutes to 1 hour by intravenous anticonvulsant medication (diazepam and phenytoin).

It is important the differential diagnosis between absence status and complex partial SE because the choice of treatment; however, this differentiation, sometimes, may be difficult. Absence status comprised all cases of nonconvulsive SE, except those in which a focality could be shown in the EEG, as in our patient. Thus, it is obvious that the clinical characteristics alone do not allow an unequivocal differentiation between these two entities. The risk of misdiagnosing a complex partial SE as absence status, therefore remains, unless an EEG of the initial phase of the status is obtained.

REFERENCES