Non-motor seizure diagnosed after mild traumatic brain injury: a case report

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INTRODUCTION

Head injury due to trauma is one of the reasons why patients are taken to the hospital. The incidence of head injury in developed countries is estimated at 1.7 million, with a mortality rate of 52,000 patients each year.¹,² Severe head injury can trigger seizures up to status epilepticus in as many as 20-30% of cases in the acute phase of head injury.³,⁴ and 52% are nonmotor seizures.⁵,⁶

Classification of head injuries based on the severity of the trauma is divided into three: 1) mild trauma, namely trauma that occurs followed by a decrease in consciousness for less than 30 minutes without a skull fracture; 2) moderate trauma, trauma accompanied by a decrease in consciousness for more than 30 minutes and less than 24 hours with/without skull fractures, and 3) severe trauma, namely trauma followed by loss of consciousness for more than 24 hours accompanied by skull fractures, hematomas, or contusions.⁷,⁸

In 10% of head injury cases, nonmotor seizures were detected by electroencephalography (EEG) performed in the acute phase.⁹ Nonmotor seizures are often not diagnosed at the start of treatment because the symptoms are often not recognized by the patient's family or doctor. The incidence of seizures in patients with minor head injuries is very low. From the literature, it was found that the type and severity of the head injury is a significant factors in the incidence of post-traumatic seizure. Thus, in this article, we report a case of nonmotor seizures in a patient after a minor head injury.

CASE DESCRIPTION

An Indonesian female, 53 years old, complained of sudden silence when speaking after 4 days earlier, she fell and hit her head on the right side. There was no loss of consciousness, headache, vomiting or limb weakness. Complaints of motor seizures and a history of previous seizures were denied. There were no cranium defects, hematomas, or open wounds on examination. The results of laboratory tests and non-contrast head CT scans were expected. On electroencephalography (EEG) examination, epileptiform waves were found in the left temporal region. The patient was given anti-seizure drugs, and complaints improved within 2 days of therapy. It is crucial to carry out an EEG examination in the acute phase of a head injury to exclude the possibility of nonmotor seizures and brain dysfunction, which will be at risk of causing seizures later in life after trauma.

Conclusion: EEG examination in the acute phase of head injury is used to exclude the possibility of nonmotor seizures and brain dysfunction, which will be at risk of causing seizures later in life after trauma.

Keywords: Brain injury, electroencephalography, a non-motor seizure.

when talking, and the patient was unaware of the incident. The family recognizes that this complaint appears up to five times per day. The patient had never had seizures or similar complaints before. A new complaint appeared shortly after the patient suffered a collision to the head when he fell. The patient has a history of hypertension and diabetes mellitus. The family admits the patient regularly takes medication to control hypertension and diabetes. There is no history of previous seizures.

On examination, the patient had a blood pressure of 180/120 mmHg, a pulse of 110 times per minute, a respiratory rate of 18 times per minute, and a temperature of 36.8 Celsius. No hematomas or open wounds were found on examination of the patient's head and neck. The patient was fully conscious, with no signs of meningeal irritation, decreased vision, visual field disturbances, or other cranial nerve abnormalities. On motor examination, there was a strength of 5 in all four extremities, and no sensory or autonomic disorders were found.

Laboratory examination of complete blood and serum electrolytes showed the expected results that were normal. Furthermore, the patient underwent a non-contrast CT scan of the head to rule out a structural brain lesion due to a collision 4 days earlier. The normal results were obtained from the head CT without contrast.

The patient's complaints persisted on the second day of treatment. The patient underwent electroencephalography (EEG) examination with suspicion of non-motor seizures. The patient's EEG results showed a Sharp wave image in the left Temporal region with a trigger factor for mental activation (Figure 1).

The patient has been given anticonvulsant therapy, Valproic acid 500 mg every 8 hours. Two days after the therapy administration, the frequency of sudden silent speech was reduced to two times per day. Monitoring at the Neurology Polyclinic after the patient finished treatment showed an improvement in the patient's complaints. After 6 months after treatment, the patient had no complaints similar to the therapy given. The evaluation of the EEG showed normal results.

**DISCUSSION**

Moderate to severe head injuries can result in seizures, which can develop over a long period. Clinicians could be tempted to believe that epileptic seizures that occur after a head injury are the cause. TBI is connected to dissociative seizures, formerly known as psychogenic non-epileptic seizures. Dissociative seizures affect women significantly three times greater rather than men. In contrast to the predominance of males who experience epileptic seizures. This might be because women are more susceptible to various risk factors. Female patients are more likely than male patients to experience post-traumatic dissociation seizures. In our case, we found a female, 53 years old, had a non-motor seizure after experiencing mild traumatic brain injury. The symptoms of nonmotor seizures are nonspecific, so the diagnosis is often missed. These symptoms include anorexia, aphasia/mutism, amnesia, lethargy, agitation/aggression, flickering, confusion, facial twitching, nystagmus, and tremors. Nonmotor status epilepticus can manifest as decreased consciousness, altered mental status, automatism, behavioral changes, and hallucinations. Nonmotor seizures are said to be an EEG diagnosis.

The previous study found that very mild trauma, in which there was no loss of consciousness and no injuries to the head or neck, did not increase the risk of seizures in the first 1 year of post-traumatic observation. However, in patients with head injuries, the risk of seizures was found to be 1.7. Another study stated that the ratio of occurrence of seizures in patients after mild head injury was only 1.5. Only Christensen et al. reported that the risk of seizures in mild head injury was doubled at 10 years of post-traumatic observation.
Structural brain lesions in neck injuries can result from an acceleration-deceleration mechanism resulting in diffuse axonal injury, venous tear, and diffuse vascular injury. Abnormalities in EEG results may not reflect an existing structural lesion but may represent transient changes due to cortical stimulation, cerebral circulation, or local post-traumatic edema. Numerous processes after injury include necrosis, micro-hemorrhage, axonal damage, apoptosis, demyelination, microgliosis, inflammation, oxidative stress, and later phases of neurodegeneration, regeneration, revascularization, and remodeling, may contribute to the circuit changes that lead to last epilepsy. In mild head trauma, seizures can also occur due to changes in the nervous tissue, including inflammatory changes and axonal growth, for some time after the trauma. However, there is still a lack of conclusive evidence regarding the causal role of neuroinflammation in epileptogenic, especially in the event of trauma.

According to a Cochrane review, studies on preventive antiepileptic medications after TBI were of low quality. They discovered that therapy (phenytoin or carbamazepine) did not affect late seizures or death after TBI but may decrease the frequency of early post-traumatic seizures. No benefit was shown when valproate and levetiracetam were compared to phenytoin. There are no recommendations in the Brain Trauma Foundation Guideline. Prophylaxis for those without seizures cannot be recommended without a better risk classification. In this case, report the patient has been given anticonvulsant therapy Valproic acid 500 mg every 8 hours. Two days after the therapy administration, the frequency of sudden silent speech was reduced to two times per day. Monitoring at the Neurology Polyclinic after the patient finished treatment showed an improvement in the patient’s complaints. After 6 months after treatment, the patient had no complaints similar to the therapy given. The evaluation of the EEG showed normal results.

CONCLUSION

EEG examination in the acute phase of head injury is used to exclude the possibility of nonmotor seizures and brain dysfunction, which will be at risk of causing seizures later in life after trauma.

DISCLOSURE

Conflict of Interest
The authors report no conflict of interest in this work.

Informed Concern
The patient was informed and agreed that the case would be published. However, the identity of the patient was kept confidential.

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Author Contribution
All of the authors contributed in all to the manuscript processes.

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