Successful Management of Paroxysmal Sympathetic Hyperactivity in Anti-N-Methyl-D-Aspartate Receptor Encephalitis

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Abstract

Background: Paroxysmal sympathetic hyperactivity (PSH) is a hyper-adrenergic clinical syndrome that is predominantly described in acquired brain injuries, less commonly in other acquired brain injuries. The association of anti-N-Methyl-D-Aspartate Receptor (NMDAR) encephalitis and PSH is not well established. Case Report: We reported a PSH associated with anti-NMDAR encephalitis and the successful management of this case. A 32-year-old female presented with episodic involuntary movement and rapidly progressed to seizure and acute confusion. Further investigations confirmed the diagnosis of ovarian teratoma associated anti-NMDAR encephalitis which was treated with pulsed methyl-prednisolone, immunotherapy, and ovarian cystectomy. She developed episodic fever, tachycardia, hypertension, diaphoresis, and dystonia during her hospital stay. PSH was suspected after other causes were excluded and the diagnosis was made more probable by applying PSH assessment measure (PSH-AM). A combination of pharmacological agents was started with good response. At 9 months from symptoms onset, she achieved almost complete recovery and returned to work. Conclusion: This case report raises the awareness of PSH in anti-NMDAR encephalitis and illustrates the use of PSH-AM as an assessment tool to diagnose PSH and assess its severity. When PSH was recognized early and treated aggressively, good outcome can be achieved.

Keywords: Anti-N-Methyl-D-Aspartate Receptor Encephalitis, Dystonia, Hypertension, Seizures, Ovarian Teratoma.

Introduction

Paroxysmal sympathetic hyperactivity (PSH) is a distinct syndrome which is characterized by paroxysmal transient hyperthermia, tachycardia, hypertension, tachypnea, excessive diaphoresis and specific posturing [1]. PSH is commonly described in traumatic brain injury (TBI) but not commonly recognized in anti-NMDAR encephalitis and the reports have been sparse. This may be contributed by the absence of a universally accepted terminology as well as standard diagnostic criteria [2]. We report a case of PSH in a patient with anti-NMDAR encephalitis and discuss the application of the recently developed assessment tool of PSH in the management of this case.

Case Report

A 32-year-old Chinese female presented to the Department of Neurology in a tertiary hospital with one day history of episodic involuntary choreiform movements and headache. The symptoms were preceded by an upper respiratory tract infection. Subsequently she developed acute confusion, agitation, and seizure. She did not have any medical or psychiatric illness, and was not on any medication or illicit drug prior to presentation. Extensive investigations were carried out including inflammatory and autoimmune markers, magnetic resonance imaging (MRI) of the brain, CT thorax, abdomen and pelvis, electroencephalography (EEG) and lumbar puncture. There was no evidence
of infection of the central nervous system (CNS) and EEG did not show epileptic form activity. MRI brain was unremarkable. However, the anti-NMDA receptor in both cerebral spinal fluid (CSF) and serum were positive. CT scan revealed a right ovarian tumor for which cystectomy was done and mature ovarian teratoma was confirmed histologically. The diagnosis of ovarian teratoma associated anti-NMDAR encephalitis was made.

On day 5 of admission, the patient was started on first line treatment including intravenous immunoglobulins 21 gm and methyl-prednisolone 1 g/day for 5 days followed by oral prednisolone in tapering dose. As her confusion and agitation still persisted 2 weeks after completion of treatment, she was considered as poor response to the first line treatment and was started on second line treatment including 3 cycles of IV cyclophosphamide 1000 mg, and 6 cycles of rituximab 500 mg from week 3 to week 10.

Two weeks after symptoms onset she developed fever, sinus tachycardia, hypertension, diaphoresis, and episodic motor posturing without obvious precipitating factors. There were 5 to 6 episodes per day with each episode lasting 15 to 20 minutes. Her highest temperature was 38.7 degree celsius, heart rate was more than 140 beats per minute, respiratory rate was more than 30 breaths per minute, systolic blood pressure was more than 180 mmHg [Fig.1]. Septic workup was not suggestive of an infective etiology. Her symptoms also did not resolve with empirical broad coverage antibiotics. PSH was suspected and PSH assessment measure (PSH-AM) [3] was applied to assess the likelihood of diagnosis. Her PSH-AM score was 26 which made the diagnosis of PSH probable [Fig.2]. The patient was started on oral clonazepam 0.5 mg every night (ON) and the dose was titrated to 1 mg three times per day (TDS) subsequently. After 1 week of clonazepam (on day 35 of admission), dystonia and diaphoresis improved significantly but fever, hypertension and tachycardia were still present episodically. PSH-AM score reduced from 26 to 18 suggesting PSH was still probable [Fig.2]. Propranolol 20 mg TDS, gabapentin 300 mg TDS were started and her symptoms resolved 1 week later. The PSH-AM assessment was repeated and the score decreased to 6 [Fig.2]. Clonazepam and gabapentin were gradually tapered down and stopped completely before discharge. Propranolol was continued at 20 mg TDS upon discharge and stopped 3 months later. There was no obvious side effect observed from the medications.

![Fig.1: Patient's respiratory rate (RR), systolic blood pressure (SBP), diastolic blood pressure (DBP), and heart rate (HR) after admission. Data point represents the daily highest record of patient's RR, SBP, DBP and HR after admission. After 2 weeks of admission, there was an increasing trend of RR, BP and HR. With starting clonazepam, there was a significant improvement in patient's RR, DBP. The RR and DBP decreased and stabilized, however SBP and HR were still on and off high. After starting gabapentin and propranolol, patient's SBP and HR normalized.](image1)

![Fig.2: Serial paroxysmal sympathetic hyperactivity-assessment measure (PSH-AM) monitor in this case.](image2)
At 3 months from symptoms onset, her neurological and psychiatric symptoms improved significantly and she was discharged well. At 9 months from symptoms onset, she achieved almost complete recovery except subtle difficulties in auditory memory. She was independent in her activities of daily living and she returned to work.

Discussion

We reported PSH in a case with anti-NMDAR encephalitis, which is a relative rare condition in the literature. Anti-NMDAR encephalitis is one of the autoimmune encephalitis syndromes which present with prodromal headache and fever, followed by a multistage progression of symptoms that include seizures, decreased level of consciousness and memory deficits. The exact incidence of anti-NMDAR encephalitis is unknown. In a study there were more than 400 cases reported in a three-year period [4]. The diagnosis of anti-NMDAR encephalitis is confirmed by the detection of IgG antibodies to the GluN1 (also known as NR1) subunit of the NMDAR in serum or CSF [5]. It has a high association with ovarian teratoma presenting as paraneoplastic syndrome [6]. Treatment for anti-NMDAR encephalitis includes tumor removal and immunotherapy. Good outcome can be expected with appropriate treatment of anti-NMDAR encephalitis. A retrospective study of 577 patients showed approximately 80 percent of patients achieved a good outcome by 24 months [6]. In our case, this patient had a typical presentation of anti-NMDAR encephalitis for which she received tumor removal, steroids and IVIG as first-line therapy. She also received second-line therapy including rituximab and cyclophosphamide. Good clinical outcome was achieved with almost complete recovery at 9 months. Recognizing and successful managing PSH has contributed to the good clinical outcome of this case.

The association of PSH with anti-NMDAR encephalitis is not well established. A review of 349 PSH case reports found that about 80% of PSH followed TBI, 10% followed anoxic brain injury, 5% followed stroke, and the remaining 5% occurred in association with hydrocephalus, tumors, hypoglycemia, infections, or unspecified causes [7]. To date there were only few case reports available describing the association of anti-NMDAR encephalitis with PSH [8,9]. Of note, although the cases formally diagnosed PSH was rare in anti-NMDAR encephalitis, autonomic instability has been reported in anti-NMDAR encephalitis with a frequency of 69-89% in various case series [10,11]. The most common manifestations of autonomic instability reported are hyperthermia, cardiac dysrhythmias (tachycardia or bradycardia), hypersalivation, hypotension, hypertension, urinary incontinence, and sexual dysfunction [10]. We believe that the autonomic instability reported in anti-NMDAR encephalitis may often be PSH and PSH can be under-diagnosed and under reported in anti-NMDAR encephalitis.

There was no clear definition or terminology to describe PSH until 2014 when an international consensus was reached with a proposal of a clinical scoring system, namely, PSH-AM [3]. The PSH-AM score is calculated by combining the clinical feature scale (CFS) score and diagnosis likelihood tool (DLT) score, which gives an estimate of the probability of a diagnosis of PSH [3]. In a case series [1], among 394 survivors of 521 patients admitted with acquired brain injury, 6 patients (1.5%) were diagnosed as PSH by using PSH-AM. This study showed that PSH-AM provides a more objective measure to increase the clinical certainty of diagnosing PSH. The diagnosis of PSH in our case was supported by PSH-AM, and patient’s progress was monitored by serial charting of PSH-AM as well. From our experience, serial collection of PSH-AM score allows clinicians to assess the severity, monitor the progress of the patient’s clinical status and response to treatment, therefore improve the efficacy in PSH management.

PSH can lead to serious complication if left untreated [2]. A wide range of pharmacological
treatment has been used to manage symptoms of PSH such as opioids, benzodiazepines, beta-adrenergic blockers, clonidine and gabapentin [2]. In practice, a combination of drugs is often needed to target the different components of PSH. The pharmacological agents chosen are largely dependent on the availability of the drugs and preference of the managing physicians as there is paucity of evidence available. In this case study, we found that propranolol, gabapentin and clonazepam were effective in controlling the symptoms of our patient. We also found that PSH-AM is a helpful measure as demonstrated in this case study as it allows us to diagnose PSH more confidently, guide the treatment and monitor the effects.

Conclusion

The diagnosis of PSH can be challenging in non-traumatic brain injuries and hence, often under-recognized. This is a case report of anti-NMDAR encephalitis complicated by PSH. This case report supported the use of PSH-AM for the diagnosis of PSH and to monitor its treatment effect. However as a single case report, it is difficult to assess the validity of PSH-AM. Larger studies are required to provide more evidence to support the use of PSH-AM and to assess whether management guided by the measure can improve outcome of the patients. Furthermore, clinical experience and judgement are important to exclude other alternative diagnoses which can mimic PSH.

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