Posterior reversible encephalopathy syndrome and acute post-streptococcal glomerulonephritis mimicking breakthrough seizures

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Abstract

We report the case of a 14-year-old boy with a past history of primary generalized seizures, who had been seizure-free for 2 years on sodium valproate and presented with generalized tonic clonic seizures suggestive of breakthrough seizures. Examination revealed hypertension, impetiginous lesions of the lower limbs, microscopic hematuria, elevated anti-streptolysin O titre and low complement levels consistent with acute post-streptococcal glomerulonephritis. Cranial magnetic resonance imaging (MRI) demonstrated changes consistent with posterior reversible encephalopathy syndrome. Hypertension was controlled with intravenous nitroglycerin followed by oral captopril and amlopidine. Brain MRI changes returned normal within 2 weeks. The nephritis went into remission within 6 months and after 8 months the patient has been seizure free again. Posterior reversible encephalopathy syndrome appeared to have neither short nor intermediate effect on seizure control in this patient. The relationship between posterior reversible encephalopathy syndrome and seizures is reviewed.

Introduction

Posterior reversible encephalopathy syndrome (PRES) associated with acute post-streptococcal glomerulonephritis (PSGN) or with other illnesses is rare.1,2 The risk of seizures following PRES and its association with diverse illnesses is a growing area of interest and a recently published longitudinal study demonstrated long term seizures in 3 out of 127 patients after a median follow-up of 3 years.3 We report the case of a boy with previ-ous seizures who presented clinically as possible breakthrough seizures so unmasking PRES and PSGN and focus on the relationship between seizures and PRES.

Case Report

A 14-year-old male student presented to Accident and Emergency Department with multiple seizures over a ten hour period. The semiology of seizures was typical with features of generalized tonic-clonic seizures with associated eye rolling, tongue biting, urinary incontinence and post-ictal drowsiness. All seizures were witnessed and the patient was conscious between seizure episodes. It was felt that these may have been breakthrough seizures. However, he had fever, generalized malaise, headaches, shortness of breath and productive cough for three days prior to admission. Facial and leg swelling was also reported over the same time period. No alteration in color or volume of urine output was reported and there was no visual disturbance. The patient was known to have a generalized tonic-clonic seizure disorder diagnosed at age 3 years, but was seizure free for two years prior to this admission and controlled on sodium valproate 600 mg orally twice daily. Previous brain imaging and electroencephalogram (EEG) were never done due to sociocultural reasons. There was no history of head trauma or drug abuse. He had normal milestones of development and lived in a rural forested area.

On examination, there was drowsiness (Glasgow Coma Scale 14/15) and some mild respiratory distress. Vital signs revealed an elevated blood pressure at 167/106 mmHg, pulse 106/min, temperature 36.9°C with a respiratory rate of 24 bpm. Dipstick urinalysis showed 4+ blood, but no protein. The chest examination revealed crepitations bi-basally and there was obvious facial and periorbital edema. There were no focal neurological deficits or signs of meningism. Plantar reflexes were upgoing bilaterally. Pupils were 3 mm and reactive to light and fundoscopy was normal. On both legs there were multiple excoriated papular lesions which were attributed to scratched mosquito bites (Figure 1).

The patient was admitted to the high dependency unit and commenced on glyceryl trinitrate. Intravenous infusion (1 mg per mL) given over 24 hours, which was titrated according to his blood pressure and intravenous phenytoin 100 mg three times daily, were used in the first 24 hours to control seizures. Once stable, oral amlopidine and captopril were started, his blood pressure was controlled on these oral agents and, after day four, he remained normotensive off all antihypertensives. Sodium valproate 200 mg orally twice daily was restarted on day 2 post admission. Sodium valproate level was unavailable.

Blood investigations showed normal renal and liver function tests. There was a low C3 level 66 mg/dL (90-180), an elevated antistreptolysin O titre 329 IU/mL (0-200) and normal C reactive protein. The erythrocyte sedimentation rate was 20 mm/hr and the white cell count was elevated at 13.3x10^9/L. Hemoglobin and platelets levels were normal. The antinuclear antibody, double stranded DNA, perinuclear anti-neutrophil cytoplasmic antibody and cytoplasm anti-neutrophil antibody were normal.

Urine microscopy revealed red blood cells: 216.3 per high power field with red cell casts. A chest X-ray was normal. Renal artery magnetic resonance angiogram, 5-hydroxyindole acetic acid, vanillylmandelic acid and metanephrine levels were all normal. Magnetic resonance imaging (MRI) changes of the brain are illustrated in Figure 2A, showing hyperintense signals bilaterally in the occipital lobes and posterior frontal/anterior parietal lobes on axial Flair image. Figure 2B shows normal T2 MRI 3 weeks later. An interictal EEG one month later was normal.

On discharge, his C3 levels had returned to normal and dipstick hematuria resolved. He was discharged on sodium valproate 200 mg orally twice daily to outpatient care. Twelve months later the patient remained seizure free.

Keywords: Posterior reversible encephalopathy syndrome; post-streptococcal glomerulonephritis; breakthrough seizures; neurology.

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Written informed consent was obtained from the patient for publication of this case report and the accompanying magnetic resonance images.

Discussion

The term PRES was coined in 1996. In that report, 15 patients were evaluated and hypertensive encephalopathy was deemed the underlying cause of this syndrome. The mechanism of the syndrome is poorly understood, but it has been postulated that a brain capillary leak syndrome, fluid retention, endothelial dysfunction/injury and hypoperfusion may be the underlying pathogenetic mechanism. Recently blood-brain barrier breakdown has been demonstrated in rats with decreased uterine perfusion pressure and it is hoped that this possible model of PRES will allow further elucidation of this illness. Our patient presented with some of the typical features of PRES, namely headaches, decreased alertness, seizures and hypertension associated with PSGN. The patient was known to have a generalized seizure disorder but had been seizure free for two years prior to presentation, demonstrating the importance of searching for new or alternative pathology in a previously well controlled seizure patient.

The relationship between seizures and PRES has aroused interest. Seizures in PRES are usually generalized and occasionally focal. Raj et al., in a study of 8 pediatric patients with PRES of diverse causes, found predominant generalized tonic and/or clonic seizures with no neurological sequel after 1 year follow up. PRES presenting as status epilepticus has been documented in a previously healthy child, two patients on chemotherapy, after pediatric hematopoietic stem cell transplantation in 10 patients and in a further 10 cases where focal-onset complex partial seizure was present in the majority of cases. PRES, in addition, can cause both convulsive and non-convulsive status epilepticus.

The EEG usually show diffuse theta/delta slowing and occasionally epileptogenic activity with focal sharp-wave and rarely periodic lateralizing epileptiform discharges and long term anticonvulsants are usually not required. Furthermore, 2 patients presenting with temporal lobe epilepsy have been found to have previous PRES, and occipital lobe epilepsy associated with atrophy of the occipital lobe and EEG abnormalities have been noted subsequent to PRES in post-partum eclampsia. Recurrent PRES in the same patient has caused chronic epilepsy. Also, hippocampal sclerosis has also been described in a recent case.

Posterior reversible encephalopathy syndrome has also been recently noted to cause long term seizures albeit in low frequency. Long term follow up however did not show seizures as a sequel in a small cohort of patients with PRES in two other studies. Thus the relationship between PRES and long term seizures is still unclear and the epileptogenic foci in PRES have not been mapped yet, most likely due to lack of an adequate experimental model. Our case is unusual since the patient had a history of previous generalized seizures unlike previous reported cases of PRES. It is likely that the PRES per se was responsible for the seizures leading to admission of this patient, but it is also theoretically possible that the seizure threshold at the epileptogenic zone was lowered by the pre-existing seizure disorder facilitating the seizures. Our observations, though only in a singular case, show that at least in the short and medium term, typical PRES did not have a detrimental effect on seizure control.

Conceivably in cases of severe PRES with ensuing permanent brain damage, the usual seizure pattern or frequency may be altered in a patient who has preexisting seizures, given the above evidence. However, we couldn’t find such reports, so this will be an area for further research. Though it appears there is full recovery in PRES in most patients with a few recorded cases of long term seizures, these reports nevertheless warn clinicians to be vigilant in diagnosis and aggressive in treatment of both hypertension and seizures in case of PRES.

Conclusions

Clinicians are therefore advised that PRES should be considered in the differential diagnosis when searching for new or underlying pathological processes during apparent breakthrough seizures; especially if predisposing factors for PRES exist and aggressive treatment should be commenced.

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