Alternative psychosis (forced normalisation) in epilepsy

V T R Ntsanwisi, MB ChB, MMed (Psych)  
S T Rataemane, MB ChB, FF Psych (SA), Child and Adolescent Psychiatry (UK)  
Department of Psychiatry, University of Limpopo (MEDUNSA campus), PO Medunsa, 0204

D S Magazi, MB ChB, MMed, FCP (Neurol) SA  
Department of Neurology, University of Limpopo (MEDUNSA campus)

Forced normalisation is a paradoxical relationship between seizure activity and behavioural problems. A 20-year-old man with recurrent refractory tonic-clonic epilepsy experienced forced normalisation while on medication with multiple anti-epileptic drugs (valproate sodium, carbamazepine and topiramate). A reduction in the seizure burden correlated with sudden behavioural changes manifesting with aggressive outbursts and violence.

The case may help clarify the mechanism of forced normalisation while providing some helpful hints regarding the diagnosis and treatment of symptoms observed in recurrent refractory seizures.

Brief episodes of abnormal behaviour have been recorded after dramatic reduction of seizures using anti-epileptic drugs (AEDs). This phenomenon is called alternative psychosis, or forced normalisation when supportive electro-encephalogram (EEG) evidence is available.

Landolt was the first to report improvement in EEG activity during periods of abnormal behaviour. The mechanism of forced normalisation is still not fully understood, although the kindling phenomenon, the phenomenon of long-term potentiation and the channel disorder paradigm have all been proposed as possible explanations. Another theory relating to this phenomenon states that epileptiform discharges may mimic electroconvulsive therapy in a local area and this seizure suppression may lead to psychopathology.

Patients with refractory temporal lobe epilepsy who undergo unilateral anterior temporal lobectomy have been observed to develop a de novo psychosis with diminished seizures. This is thought to be an alternative psychosis related to forced normalisation of the EEG.

The absence of clear diagnostic criteria for forced normalisation has been an impediment both to its routine diagnosis and to further research. Research into this phenomenon has been confined mainly to case reports (as in this case) or to retrospective studies.

Anticonvulsant drugs have been associated with new-onset depression and psychosis. If anticonvulsants have recently been changed, this should always be considered as a potential cause of a new or worsening depressive or psychotic illness.

There are case reports in the literature describing psychosis relating to topiramate, zonisamide and levetiracetam (see Table I for a list of drugs implicated in forced normalisation). Some of these reports may relate to the process of forced normalisation, in which a diminished frequency of seizures allows psychotic symptoms to emerge.

Krishnamoorthy and Trimble list questions posed by other researchers that still remain unanswered:

1. Should the EEG necessarily be completely normal for this diagnosis to be made, or should relative normalisation also be included?
2. What is the relationship of the EEG to suppression of seizures?
3. Is alternative psychosis the expression of forced normalisation, a variant, or unrelated to it?

The current approach is wider inclusion of cases that show a decrease in seizure frequency with both relative and complete normalisation of the EEG.

The following are the proposed criteria for forced normalisation:

Primary (essential) criteria:

1. Established diagnosis of epilepsy based on clinical history, EEG and imaging
2. Presence of a behavioural disturbance of acute/sub-acute onset characterised by one of the following:
   - psychosis with thought disorder, delusions, hallucinations
   - significant mood change, mania/hypomania or depression
   - anxiety with depersonalisation, derealisation
   - hysteria: motor, sensory, abasia

Patients with refractory temporal lobe epilepsy who undergo unilateral anterior temporal lobectomy have been observed to develop a de novo psychosis with diminished seizures.
3A. Reduction in the total number of spikes counted in a 60-minute awake EEG recording with a normal 16-channel machine, using standard 10-20 electrode placement, by over 50% compared with a similar recording performed during a normal state of behaviour or
3B. Report of complete cessation of seizures for at least 1 week, corroborated by a relative or a carer.

Supportive criteria:
1. Recent change (within 30 days) of pharmaco-therapeutic regimen
2. Report of similar episodes of seizure cessation and behavioural disturbance in the past, from close relative, carer or general practitioner, or documentation of these in hospital records, with or without EEG evidence. This may or may not be linked with anticonvulsant drugs.

The diagnosis is made in the presence of:
Primary criteria 1, 2 and 3A
or
Primary criteria 1, 2 and 3B and one supportive criterion. 

Trimble makes no specific recommendation regarding the type of imaging of epilepsy; however, according to Kanner et al., the presence of interictal psychosis should prompt the clinician to order high-resolution magnetic resonance imaging (MRI) in search of small tumours (hamartomas and ganglionomas), some of which may not be detectable on the standard MRI and may cause refractory epilepsy. Fluorodeoxyglucose-positron emission tomography (FDG-PET) may show hypometabolic areas that correlate with epileptic foci as revealed by the EEG. However, the challenge in the South African context is limited availability of some of these diagnostic tools.

Case report
A 20-year-old man with recurrent refractory tonic-clonic epilepsy was referred to the liaison psychiatry registrar at George Mukhari Hospital by the neurology registrar after he had an altercation with the nursing staff in the neurology ward.

The patient had been treated for recurrent refractory epilepsy by the neurology department since 1996. He was on multiple AEDs. His current drug regimen was sodium valproate 1 000 mg in the morning and 1 500 mg at night, carbamazepine 400 mg twice a day and topiramate 50 mg twice a day. There had been a substantial reduction in his seizure burden from a seizure approximately every 2 days a year ago before the introduction of the current drug regimen to 3-4 seizures a month after its introduction. The change in behaviour had been observed for the first time by the family in the 6 months following the reduction in seizure frequency.

For the first time, the patient had presented with bouts of verbal and physical aggression and even violence. He had recently assaulted his grandmother and also chased his elder brother with a knife with the intention of harming him. According to the family there had been no provocation prior to the incident, and the patient subsequently apologised, showing remorse upon realisation of the wrongfulness of his actions. The patient admitted to taking alcohol occasionally at social functions. He drank only beer, three or four 350 ml cans at a sitting. He denied smoking cannabis or taking any other illicit substance. There was a positive family history of epilepsy, his deceased maternal aunt having been affected.

According to his mother, the patient had been delivered vaginally at term, although the labour had been prolonged and he had...
a 2-week stay in the neonatal ward due to severe neonatal jaundice. The developmental milestones were reported to be normal. According to the mother the patient had passed every class at the first attempt up to Grade 5 in 1999, but was forced to stop school because of his severe epilepsy. However, no school report was available to substantiate her account of his academic performance. His premorbid personality was described as quiet and soft-spoken. According to his mother, the patient had undergone a personality change in the past 12 months, having become impulsive, aggressive and violent. He became prone to temper tantrums, banging doors, and would throw himself to the floor if he did not get his way.

The findings on physical examination were normal, with a blood pressure of 130/90 mmHg, a regular pulse rate of 72 beats/min, and no evidence of lymphadenopathy or thyroid enlargement. The cardiovascular, respiratory, musculoskeletal and abdominal systems appeared normal. The pupils were equal and reactive to light. The optic disc appeared normal on fundoscopy. The cranial nerves were intact and examination of the upper and lower limbs revealed normal power, tone, reflexes, co-ordination and sensation.

On mental status examination, the patient’s behaviour was appropriate, though conflict was observed in his interaction with his mother, leading to irritable mood. Affect was appropriate, and there was mild psychomotor retardation. His speech was slow in rhythm but normal in quantity and volume. He was orientated to person, place and time. Attention and concentration were poor; he was able to do the first two of the serial 7 subtraction (93, 86), but there was latency of response, pointing towards slowing of cognitive speed. Memory was intact, and there were no perceptual distortions.

The patient’s thinking process was logical and goal directed, and the content showed no psychopathology. Insight was partial and judgement was fair, as evidenced by the retaliation after provocation, and then showing remorse. Abstract thinking was normal. His intelligence was borderline towards sub-average. He had a Mini-Mental State score of 23/30, pointing towards cognitive deficit.

Since there was no aura, the patient was clinically classified as having primary generalised epilepsy. The EEG showed multifocal spike and wave abnormalities.

This patient’s clinical course meets the criteria for alternative psychosis and forced normalisation of the EEG. We accommodate the possibility of coincidence, in other words no causal link between seizure frequency and the emergence of aggressive behaviour. In this case, however, we are persuaded towards there being a causal link in that this was a new development that emerged in the 12th year only after a change in drug regimen that resulted in a drastic reduction in seizure frequency. The patient satisfies Krishnamoorthy and Trimble’s criteria 1, 2, 3B and one supportive criterion.

A differential diagnosis of episodic dyscontrol syndrome was made, this is characterised by sudden episodes of spontaneously released violence with minimal provocation that terminate abruptly. These episodes are precipitated by small amounts of alcohol and anticonvulsant drugs, especially the benzodiazepines, and are considered to be a form of paradoxical reaction that occurs in individuals with pre-existing brain damage, such as represented by this patient, who sustained neuronal injury as a result of recurrent refractory seizures.21

Psychosis secondary to epilepsy is an important consideration in this case scenario. Patients with chronic epilepsy (especially those with temporal lobe epilepsy) eventually succumb to a schizophrenia-like state.22

Discussion

The patient had been suffering from recurrent epilepsy since 1996, when he was 8 years old. His medical records show that he was initially treated with phenytoin, which was stopped due to toxicity and gingival hypertrophy.

Different regimens were tried without much success. Recently his seizure burden reduced to 3 - 4 seizures a month from a seizure every 2 days, but then episodes of abnormal behaviour characterised by aggressive outbursts and impulsivity came to the fore for the first time (alternative psychosis or forced normalisation). The new adjunct AEDs have been observed to be associated with this phenomenon. Trimble and Schmitz22 have warned clinicians about these side-effects and have suggested slower titration of medication for seizure control as a measure to prevent them.

Initial drug therapy to control seizure activity is based on the specific type of seizure. Monotherapy is initiated with a single agent until seizure remission or signs of toxicity. Patients on monotherapy fare much better than those on combination therapy regarding compliance and side-effects. In the event of failure to control seizures with the first drug, monotherapy with a different antiepileptic is suggested before resorting to augmentation strategies with different AEDs.23,24

Most AEDs exert their effect by inhibiting generation of an electric discharge from a focal area or by blocking the propagation of an abnormal electric charge to an adjacent brain area. They accomplish this by a variety of mechanisms, including:

- inhibition of voltage-gated channels (Na+, K+ or Ca2+)
- enhancement of inhibitory GABAergic impulses
- attenuation of excitatory glutamatergic transmission 25,26,27
Systemic data on the psychiatric adverse effects of primary AEDs are very limited and mainly based on empirical data and small case series (Table III).

Drugs with a GABAergic mechanism of action have sedating, anxiolytic and anti-manic properties, while drugs with a glutamatergic mechanism of action have activating, anxiogenic and antidepressant effects. Tiagabine and vigabatrin are examples of the first category, and lamotrigine an example of the second.

Topiramate was the second-line augmenting drug used in the present case. Its manner of action is multimodal and involves blockage of voltage-gated sodium channels, potentiation of GABAergic transmission and inhibition of excitatory glutameric pathways through an action at the AMPA receptor site.24

Neuropsychiatric manifestations in the form of psychosis, affective disorders and behavioural disorders have emerged in patients on topiramate, vigabatrin and tiagabine. Trimble and Rusch25 carried out a retrospective case study on 89 patients who developed behavioural symptoms during treatment with topiramate, vigabatrin and tiagabine. It was found that 99% of the patients suffered from complex partial seizures with or without secondary generalisation. More than half were on polypharmacy with two or more AEDs. Nearly two-thirds had a previous psychiatric history.

A strong association was found between the type of previous psychiatric illness and the type of emerging psychiatric problem. A seizure-free period was observed in more than half of the patients before they developed the psychiatric symptoms,29 entrenching the perceived paradoxical relationship between reduction in seizure frequency and emergence of psychiatric illness and the type of emerging psychiatric problem.29

The goal in the treatment of epilepsy is to render the patient seizure-free and to maintain adequate control of mood and anxiety disorders after temporal lobectomy.26

Topiramate was the second-line augmenting drug used in the present case series (Table II).

Finally, we suggest a change in the nomenclature, as the current terms ‘forced normalisation’ and ‘alternative psychosis’ do not adequately describe the clinical cause and outcome of this condition. We propose the new name of ‘ictal suppression disorder’, which aptly describes this phenomenon and removes the ambiguity created by the current terms.

References