Clinical Reasoning: A 25-year-old woman with recurrent episodes of collapse and loss of consciousness

Section 1

A 25-year-old woman presented to the neurology clinic describing 3 episodes of transient loss of consciousness over 9 months. The first 2 were unwitnessed, with no prodromal symptoms. The third event was a witnessed collapse while shopping. It was reported that both the patient’s arms were stiff and by her sides with accompanied lateral tongue biting, a blue discoloration to her face, choking noises, and dilated pupils. The event lasted for less than 2 minutes, after which she was confused and drowsy. She remembered a premonitory feeling before the third collapse but found this difficult to describe.

The patient’s medical history included migraine and mild depression; she took amitriptyline 20 mg daily and citalopram 20 mg daily. She had been born at 26 weeks’ gestation but had no medical complaints or adverse early-life events, had normal developmental milestones, and was a university student. There was no family history of note. On direct questioning she described symptoms of infrequent but potent déjà-vu as well as general fatigue and lack of focus. She smoked 8–10 cigarettes per day and drank a small amount of alcohol on social occasions. She had a balanced diet and was a meat-eater.

On examination, power and sensation were normal throughout. Bilateral optic disc swelling was seen on fundoscopy; the remainder of the cranial nerve examination was normal. The patient had globally brisk reflexes, more prominent on the right (left 3+, right 4+). She had increased tone in her lower limbs, right greater than left, but plantar responses were downgoing. Cardiorespiratory and abdominal examination was normal and she had no neurocutaneous markers.

Questions for consideration:
1. What is the differential diagnosis for these events?
2. What investigations would you consider?
Section 2

The differential for unwitnessed collapse and loss of consciousness ranges from the common such as epileptic or provoked seizures, syncope, and dissociative attacks to rarities including insulinoma or colloid cyst of the fourth ventricle. The most common differentials for episodes of loss of consciousness are seizures, both spontaneous and provoked; reflex vasovagal syncope; and cardiac syncope. Thus ECG is essential in all cases of transient loss of consciousness. Recurring events narrow the differential by facilitating identification of triggers or sequences of events and increasing the likelihood of an event being witnessed. The stereotypy of recurrent events is an essential feature of seizures. A witness account is helpful but not infallible and limb jerking secondary to transient hypoxia may be misinterpreted as tonic-clonic seizure activity. Questionnaires such as that developed by Sheldon et al. can facilitate differentiation of seizures from syncope.

Physical examination can be supportive if lateralizing signs are identified and should include visual field assessment for identification of structural lesions within the optical pathway. General examination may also yield findings such as skin changes seen in tuberous sclerosis or neurofibromatosis. Further clues as to cardiac or syncopal causes include findings such as irregular pulse or postural hypotension. Our patient had a normal cardiorespiratory examination and an ECG was performed that was unremarkable, making these less likely.

In this case, the presence of difficult to describe aura, lateral tongue bite, and a prolonged postictal period favored an epilepsy diagnosis. The witnessed tonic phase and abnormal physical examination are also supportive of a structural lesion in the left hemisphere and thus in this case a clinical diagnosis of probable focal epilepsy was made. The American Epilepsy Society guidelines recommend EEG and brain imaging for adults with a suspected unprovoked seizure. EEG and brain MRI were performed (figure, A and B) to support the clinical diagnosis and classification of the seizures, to enable prognostication, and

Figure Illustrative section of EEG displayed on a double banana montage and axial T2 MRI sequences

(A) Double banana montage. Thick vertical lines demarcate 1 second of data. The highlighted area shows a high-amplitude, sharpened alpha rhythm over the right posterior temporal/occipital region. (B) Axial T2 MRI scan shows multifocal lesions with calcification (marked by arrows) indicative of a chronic process. The images on the far right are T1-weighted pre and post (upper right and lower right, respectively) gadolinium contrast administration and identify a ring-enhancing lesion. There are diffuse T2 changes (marked with arrowheads) within both the cerebral and cerebellar hemispheres. Perfusion imaging (not illustrated) showed increased perfusion within the enhancing lesions and diffusion-weighted imaging (not illustrated) showed no evidence of restricted diffusion.
to investigate for abnormal electrical activity supportive of seizures and structural brain lesions, respectively.

The EEG demonstrated a well-formed background with a sustained and responsive 10–11 Hz alpha rhythm but with marked asymmetry; at times the alpha rhythm recorded over the right posterior temporal/occipital region appeared high-amplitude and sharper in outline (figure, A). No interictal electrographic seizure activity was identified and the lead I ECG recorded simultaneously showed sinus rhythm.

MRI brain showed multifocal lesions with calcification, indicative of a chronic process, and gadolinium enhancement and diffuse T2 white matter changes within both the cerebral and cerebellar hemispheres (figure, B).

**Questions for consideration:**
1. What is the differential diagnosis of the etiology of the seizures based on the additional results?
2. What further testing should be arranged based on these MRI findings?
Section 3

The most likely diagnosis for the imaging changes, including ring enhancement, at this point is an atypical cerebral infection: the primary radiologic diagnosis, in the absence of systemic disease, was neurocysticercosis. The presence of gadolinium ring enhancement suggests an active process. It was important to promptly establish evidence for neurocysticercosis because of treatment implications.

Neurocysticercosis is a common cause of acquired epilepsy worldwide.4 Investigations should include routine blood tests, cranial imaging, ophthalmologic review, plain X-rays of muscles to screen for peripheral cysts, and stool examination for intestinal taeniasis. CSF may be tested using ELISA. Medical management is both symptomatic (e.g., antiepileptic agents for seizures) and specifically directed against the parasite (for example, albendazole) depending on the clinical state.5 Surgical intervention may be required in instances of hydrocephalus or intractable seizures with calcified cysts. CT has a role in identifying cysts with variable degrees of calcification.

Imaging features of neurocysticercosis depend on the stage of the disease. At the larval stage, the scolices may produce small (5–20 mm) lesions with little mass effect and no edema. Later, the cysts are larger, and the scolex may be visible within the cyst. Parenchymal cysts are described as vesicular, colloidal-vesicular, granular-nodular, and nodular-calciﬁed as neurocysticercosis progresses. At the later, calcified stage of the disease, the scolices are usually no longer visible.6 The use of MRI modalities may aid diagnosis, with fluid-attenuated inversion recovery (FLAIR) being more sensitive to intraventricular cysts and susceptibility-weighted imaging better for calcified lesions.6

We identified no risk factors for neurocysticercosis or atypical infection. Routine blood tests showed a raised white cell count (12.4) and C-reactive protein less than 5. Further inquiries revealed no history of foreign travel or other clear source of exposure to taenia parasites. The patient had no contact with tuberculosis, and previously had the Bacillus Calmette-Guerin vaccine. She had chicken pox as a child and had not subsequently had shingles. A viral screen for cytomegalovirus, Epstein-Barr virus, HIV, hepatitis C, and hepatitis B was negative.

CSF sampling showed no white cells, protein and glucose counts within normal limits, and negative cysticercosis ELISA. Fecal samples showed no ova, cysts, or parasites. CT thorax, abdomen, and pelvis and X-ray imaging of the thigh showed no peripheral cysts or calcifications.

**Question for consideration:**

1. Based on the negative test results, how would you expand the differential?
Section 4

The differential diagnosis for cysts with calcification includes cryptococcus, echinococcus, toxoplasma, and tuberculosis. Multiple abscesses would be unlikely with calcification, no fever, and no clear source of infection. Tuberculosis was also believed to be unlikely with no other area of abnormality and normal imaging elsewhere. Endemic fungi seldom calcify in the active phase and the patient had no history of travel to endemic areas. No positive confirmatory findings were made, leaving us without a history of exposure to cysticercosis and no significant supporting diagnostic features of neurocysticercosis (see section 3). Excision biopsy may be considered for cases of diagnostic uncertainty. In this instance, the case was discussed with colleagues (H.K. and S.T.) from an international center where neurocysticercosis is commonly encountered and with greater expertise in identifying neurocysticercosis. The lesions and extensive white matter changes were believed to be atypical for neurocysticercosis and there were no visible scolices.

Once intracerebral infection had been excluded, the differential diagnosis of leukoencephalopathy with calcification, which includes conditions such as Aicardi-Goutières syndrome and adult-onset leukoencephalopathy with axonal spheroids and pigmented glia, became the focus of investigation. Leukoencephalopathy with calcification and cysts (LCC) was deemed the top differential of these because of slow progression and juxtaposition between widespread radiologic features and modest clinical signs.

Genetic testing performed at the University of Manchester, UK, identified sequence variants in the SNORD118 gene by direct sequencing. The patient was found to be heterozygous for the n.72A>G sequence variant and the n."9C>T sequence variant. Both variants are associated with LCC and their presence was confirmed in the National Health Service laboratory. Genetic counseling was provided. In 2016, the genetic basis of LCC was identified, molecularly differentiating this disease from Coats plus due to biallelic mutations in components of CST, a key structural protein complex. LCC and Coats plus demonstrate a remarkably similar neuro-radiologic picture, the clinical difference being the presence of non-neurologic features in Coats plus.

Our patient’s seizure control improved and she remains on a single antiepileptic drug (AED). One year later, her seizures are rare and her mood is much improved.

Discussion

This case exemplifies how a rare condition may present with a familiar clinical picture; in this case, the patient’s initial presentation was not unusual in terms of recurrent collapse, with at least one convincingly described seizure. Semiology suggested a focal epilepsy and asymmetrical examination findings in conjunction with her EEG strongly suggested a structural abnormality.

Abnormalities in EEG or cranial imaging warrant further investigation and suggest an increased likelihood of seizure recurrence, and therefore the utility of AEDs. The American Academy of Neurology recommends that patients presenting with a first unprovoked seizure who have a subsequent seizure should have AED treatment offered to them because the risk of more seizures is high: 57% by 1 year and 73% by 4 years.

LCC is a rare autosomal recessive disorder characterized by variable neurologic decline with spasticity, seizures, and a movement disorder. It is a cerebral microangiopathy related to, but distinct from, Coats plus. The largest series to date shows the age at presentation varies from under 6 months to 54 years, with symptom onset at school age being most common. A total of 14 of 40 patients first presented with seizures. The demonstrated prognosis was variable, with 35 of 40 patients still living and the oldest being aged 54 years at time of publication.

A recent case study of an 18-year-old patient treated for 1 year with bevacizumab, a vascular endothelial growth factor inhibitor, showed improvement in bradykinesia and range of movement and cessation of deterioration of speech, gait, and dystonia.10 This correlated with a reduction in cyst load on MRI and reduced FLAIR-demonstrated white matter involvement.

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Disclosure

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Appendix Authors

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<thead>
<tr>
<th>Name</th>
<th>Location</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jack Wildman, MBBS</td>
<td>Royal Victoria Infirmary, Newcastle-Upon-Tyne, UK</td>
<td>Writing of the manuscript</td>
</tr>
<tr>
<td>Mark R. Baker, PhD, FRCP</td>
<td>Royal Victoria Infirmary, Newcastle-Upon-Tyne; Institute of Neuroscience, Newcastle University, UK</td>
<td>Clinical insight, revision of text</td>
</tr>
<tr>
<td>D. Ashley Price, MD, FRCP</td>
<td>Royal Victoria Infirmary, Newcastle-Upon-Tyne, UK</td>
<td>Clinical insight</td>
</tr>
<tr>
<td>Sarbesh Tiwari, MD</td>
<td>Institute of Neurosciences Kolkata, India</td>
<td>Clinical insight</td>
</tr>
<tr>
<td>Hirshikesh Kumar, MD</td>
<td>Institute of Neurosciences Kolkata, India</td>
<td>Clinical insight</td>
</tr>
</tbody>
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**References**


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**Appendix**

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<th>Name</th>
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<th>Contribution</th>
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</thead>
<tbody>
<tr>
<td>Gillian I. Rice</td>
<td>Faculty of Biology, Medicine and Health, University of Manchester, UK</td>
<td>Genetic analysis</td>
</tr>
<tr>
<td>Yanick Crow, PhD, MRCP</td>
<td>MRC Institute of Genetics and Molecular Medicine, University of Edinburgh, UK; Institut Imagine, Descartes University, Paris, France</td>
<td>Genetic analysis</td>
</tr>
<tr>
<td>Rhys H. Thomas, PhD, FRCP</td>
<td>Royal Victoria Infirmary, Newcastle-Upon-Tyne; Institute of Neuroscience, Newcastle University, UK</td>
<td>Lead clinician, revision of text, guarantor</td>
</tr>
</tbody>
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**References**


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