A 35-year-old woman with a history of substance abuse and a seizure disorder presented to the emergency department with altered mental status. According to her family, she was behaving oddly for the past several days, including standing and staring throughout the day, not speaking and resisting taking medications. The family had not witnessed this behaviour in the past. They did endorse a substance abuse history, including opiates, for which she had previously been on methadone. On examination, the patient was persistently hypertensive and tachycardic, aroused to voice, but was otherwise nonverbal. She followed minimal simple commands and had diffusely slowed movements. An MRI brain was performed on day 1 (figure 1A) that was unremarkable with the exception of a non-specific T2 hyperintensity on the L posterior centrum semiovale of unclear significance. She had no prior MR imaging for comparison. Prolonged electroencephalogram captured a right posterior quadrant seizure without clear clinical correlate, and at this time it was presumed that her abnormal behaviour was secondary to intermittent seizure activity. The patient’s home levetiracetam dose was increased from 1000 mg twice a day to 1500 mg twice a day. However, the following day, the patient then developed arrhythmic, distractible whole body shaking spells without electrographic correlate suggesting a mixed epileptic and non-epileptic disorder. On days 4–5 since admission, her examination further deteriorated, now with spasticity in the upper extremities, a left Babinski sign, and a retarded catatonic-like state without response to noxious stimuli. A repeat MRI brain obtained at this time (figure 1B) showed interval development of diffuse T2 white matter hyperintensities throughout both cerebral hemispheres with patchy regions of gyriform thickening and oedema. Additionally, there was interval worsening of diffusion restriction and multifocal petechial haemorrhage throughout the cortex of both cerebral hemispheres (figure 2). This was concerning for a toxic leukoencephalopathy most likely secondary to substance abuse.1 The patient was started on empiric antioxidants (Coenzyme Q10, Vitamin E, Vitamin C) due to possible mechanism of mitochondrial dysfunction. Concurrently, the patient was treated briefly with empiric acyclovir, while the infectious workup was pending and ultimately

Learning points

- Toxic leukoencephalopathy can occur from snorting oxycodone with delayed onset and continued progression, despite discontinuation of the substance.
- Clinical presentation depends on the degree of white matter disease and consists of abnormal vital signs with tachycardia and hypertension, neurobehavioural deficits that include inattention, pseudobulbar affect and apathy before development of tremors and spasticity on examination.
- Our findings suggest that the classic neuroimaging findings may lag behind clinical symptoms, and that the changes seen on MRI rapidly change over the course of 5–10 days.
Images in... returned negative. An inflammatory or demyelinating process was also considered, however, there was no additional evidence on serum and cerebrospinal fluid studies or MRIs of the cervical spine to support this. Over the next week, the patient gradually began speaking in stunted sentences with a flat affect and confirmed snorting crushed oxycodone and benzodiazepine tablets 3 weeks prior to presentation. She denied intravenous drug use or inhaling heroin vapours. While toxic leukoencephalopathy from substance abuse is more commonly associated with inhaling heroin vapours, also called ‘chasing the dragon’, it has also been seen with use of other opioids and administration methods. Finally, a third MRI of the brain obtained 11 days after presentation showed interval improvement of the diffuse T2 hyperintensities correlating with her improving examination. The following images demonstrate the progressive evolution of the patient’s brain MRI, consistent with toxic leukoencephalopathy most likely secondary to snorting crushed opiates.

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