Clinical Toxicology

Strychnine poisoning as an unusual cause of convulsions

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Summary: A fatal case of strychnine poisoning is presented. The patient vomited then suffered a series of tonic convulsions which were triggered by tactile stimulation. In between paroxysms he was initially alert. Eventually the patient became comatose due to anoxia and had a cardiac arrest. He presented with a marked metabolic acidosis and rapidly developed renal failure caused by acute rhabdomyolysis. This clinical picture is classical for strychnine poisoning and the complications which the intoxication produces. Attention is drawn to the fact that survival can even follow the ingestion of very large doses of strychnine providing there is no delay in diagnosis and treatment.

Introduction

Strychnine ingestion is an uncommon and a peculiarly unpleasant form of poisoning.¹ The analeptic nature and mechanism of action of the alkaloid has long been recognized.² Although low in the differential of causes of convulsions the classical clinical picture of strychnine poisoning is important to recognize because with early and appropriate treatment survival can follow the ingestion of even very large doses.³ We present the case of a farmer who died from strychnine poisoning. Although we cannot deduce from our case that early diagnosis and intervention may be life-saving, this has been well documented.⁴ We feel, however, that our case graphically illustrates the presentation of strychnine poisoning.

Case report

A 32 year old farmer was admitted to this hospital in acute renal failure of unknown aetiology. The history, obtained from his wife and ambulance crew, was that he had woken his wife at 05.00 h telling her that their nearby barn was on fire. The couple removed farm equipment from the barn but his wife later denied any possibility of smoke or fume inhalation. Thirty minutes later, while waiting for the fire brigade, the farmer vomited and after a few minutes had a series of tonic convulsions. Between convulsions he was alert and calling for help, but any physical contact appeared to cause pain, panic and further convulsions. After several paroxysms of convulsions he became cyanosed and lost consciousness. On arrival of the fire brigade and ambulance services no pulses were palpable and cardiopulmonary resuscitation was given until an output was restored. On arrival at the local hospital he was no longer cyanosed and appeared well perfused with a sinus tachycardia. Further convulsions were controlled with intravenous diazepam, paralysis and mechanical ventilation. Initial investigations performed some 15 minutes after restoration of a cardiac output, and as the patient was being intubated, revealed metabolic acidosis (pH 6.6; Pco₂ = 7.91 kPa, Po₂ = 38.2 kPa on 100% oxygen by mask) and a plasma creatinine of 130 µmol/l. Despite the correction of the acidosis with 200 ml of 8.4% intravenous sodium bicarbonate and maintenance of an adequate blood pressure (systolic blood pressure > 130 mmHg) the urine output remained less than 15 ml/h. He was transferred to this hospital for dialysis, with a presumptive diagnosis of carbon monoxide poisoning.

Investigations on admission: plasma creatinine = 410 µmol/l; creatine kinase > 1500 units per litre (normal less than 175 units per litre); carboxyhaemoglobin level 2%.

Microscopic crystals found in the urine raised the question of oxaluria secondary to ethylene glycol poisoning – for which treatment was instituted. However, gas chromatography, performed on a 1 foot column packed with 0.8% THEED on carbopack C at 110°C, indicated that ethylene glycol was not present in the plasma. Furthermore, X-ray diffraction studies revealed that the crystals were not oxalate – but remained unidentified. In view of this, further toxicological investigations were carried out on the urine. In addition to a benzodiazepine presumably administered after admission – thin layer chromato-
stimulation is sensory when dissolved Strychnine and cocaine. When dissolved in the body, Strychnine and cocaine exert an excitatory action on the medulla and, curiously, enhances the sensations of touch, smell, hearing and sight. Other reports of Strychnine poisonings have noted vomiting, hyperthermia and horizontal pendular nystagmus as clinical manifestations. Our case was thus typical, with vomiting and hypersusceptibility to external stimuli so that any attempts to help the farmer caused pain and further convulsions.

Strychnine is readily absorbed after oral administration. It is thought to be metabolized mainly in the liver, with a plasma half-life of 10 hours although up to 20% of a dose is excreted unchanged in the urine.

Fatal doses of Strychnine are reported to be as low as 5–10 mg but, more significantly, survival can follow ingestion of very high doses (over 3500 mg). If recognized in time, symptomatic treatment with diazepam or barbiturates to control the convulsions and paralysis and mechanical ventilation where necessary can prevent death from anoxia or from the associated complications of lactic acidosis, rhabdomyolysis and acute renal failure. In our case, acute renal failure developed rapidly and was presumed to be secondary to rhabdomyolysis. Death, however, was primarily due to anoxic brain damage.

We conclude that Strychnine ingestion should be considered in all cases of convulsion of obscure aetiology, especially if the person is an agricultural worker or is known to have access to the poison. If available, a carefully taken history from a witness may help make the diagnosis because of the classical clinical picture of Strychnine poisoning. Prompt and adequate anticonvulsant treatment followed by effective artificial ventilation can be life-saving.

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References