Dear editors,

A 42-year-old man was found at a bus stop in the evening. He had had a seizure and impaired consciousness. The emergency service administrated diazepam (10 mg) and midazolam (5 mg) intravenously and supported his vital functions. He was admitted to a neurological ICU at a hospital. Upon admission, he was in a deep coma with slow photoreaction, absence of other brain reflexes and no meningeal signs. Long-lasting algetic impulses indicated symmetrical flexor responses of the limbs. Oxygen saturation was 99% with an oxygen mask; blood pressure was 148/96; heart frequency was regular and slow with 43–7/min. A CT scan of the brain and angiography of the cerebral arteries showed formed hypodensity only at the right lower temporal region. This was most likely posttraumatic. An X-ray of the cervical spine was without any change in his neurological status.

His history was unremarkable. He was a non-smoker and non-drinker. No drug allergy was known. He had been treated with naproxen and paracetamol. He had no previous medical history and had no neurological or mental illness. He had been previously admitted to hospital for an assault and was treated with 0.5 mg of atropine intravenously without any change in his neurological status. He was in a deep coma without any change in his neurological status. Upon admission, he was in a deep coma with slow photoreaction, absence of other brain reflexes and no meningeal signs.

The patient was transferred from the ICU to a standard department, where he complained of moderate headache, drowsiness and slight dysarthria only. After 7 days of hospitalization, he was released without any medication.

A note on the history of our patient: 2 days before the onset of intoxication he was released from a prison. He traveled to the city to settle some formalities. He took baclofen and clonazepam orally for pain of the cervical spine and shoulders. He had suffered from minor bone injuries. Fifteen years ago, he had been briefly hospitalized for toxic hepatopathy. Since 2010, sporadic primary generalized epileptic paroxysms had been reported for which he first received phenytoin and later clonazepam (2 x 2 mg).

**Discussion**

Baclofen is a centrally acting, lipophilic derivative of gamma-aminobutyric acid. It binds to presynaptic GABA-B receptors within the brainstem and dorsal horn of the spinal cord. It acts by restricting calcium influx into the presynaptic nerve terminal, thereby reducing release of presynaptic transmitters [1]. Baclofen is used in the treatment of muscle spasticity in patients with spinal or cerebral disorders [2]. Baclofen overdose may occur in healthy individuals who have ingested baclofen intentionally. Baclofen has also been promoted as a “fun drug” by several anecdotal experiences published on the internet [2]. Our patient took baclofen and clonazepam due to pain and muscle spasms in the vertebral cervical-thoracic region but the ingested dose was not known. The drug also appeared to be used by the patient recreationally.

Adverse events of baclofen include drowsiness, headache, dizziness, and occasionally, orthostatic hypotension. Incidental or accidental overdose of this drug can cause profound central nervous system depression, including coma, hypotonia, respiratory...

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Depression, seizures and cardiovascular effects such as bradycardia [3]. Coma with restricted brain functions frequently demands artificial ventilation. Most cases feature bradycardia (35–40 min) with only mildly reduced to normal blood pressure, without elevated intracranial pressure on the brain CT or MRI. Our patient was found at the bus stop in a coma with convulsions. At the ICU admission, his bradycardia was regular with 43 min. His blood pressure and respiratory rate were also normal. Oxygen saturation was sufficient (99% with oxygen mask) and this is why the patient was not intubated.

After oral ingestion, baclofen is rapidly absorbed from the gastrointestinal tract. Blood levels peak at 2 hours. The serum half-life is 2–6 hours but can be significantly prolonged after overdose. The majority of the drug is excreted unchanged in urine. Signs of toxicity have been reported after ingestion of as little as 100 mg of baclofen [4]. 180–800 mg [5] or 100–1500 mg [6] represents a dosage that poisons the body when ingested. Coma in severe baclofen intoxication lasted at least 100–110 hours and most patients regained consciousness quickly [5]. They were discharged within a few days [6]. However, baclofen intoxication may also result in a very prolonged and profound coma and may in fact mimic brain death [7]. Our patient regained consciousness over the course of a few minutes after a 12-hour coma. Flumazenil was used in patients who had co-ingested benzodiazepines in order to reverse the comatose state [5]. Our patient co-ingested clonazepam and thus flumazenil and naloxone were injected but without any change in the deep coma. Baclofen blood level was extremely high at 2700 ng/ml but the patient’s ventilation was sufficient and only an oxygen mask was used with good results. Recovery from coma was fast and occurred within few minutes. The duration of coma was 12 hours only and did not require artificial ventilation, though the baclofen blood level was high. We assume that long-term alcohol abuse and misuse of some drugs could explain the patient’s ability to manage this high dose of baclofen so easily. He had been released from prison 2 days prior to his hospitalization and this may explain his perfect health and fast recovery.

Severe overdose of baclofen can lead to muscle flaccidity, coma and seizures. The proconvulsive effect of baclofen is believed to be the result of an activation of postsynaptic GABA-B receptors that have a depressant effect on inhibitory interneurons, leading to epileptogenesis [2]. An EEG can be helpful in differential diagnosis, showing typical abnormalities seen in a few other forms of intoxication only. The usual finding during intrathecal baclofen overdose is a reversible increase of slow wave activity, with periodic shape waves associated with periodic epileptic discharges [8]. Other abnormalities such as triphasic waves and burst suppressions have been described in cases of oral overdose [1,9]. Our patient was found at a bus stop in generalized convulsion and coma. The EEG was abnormal with a marked slowing, bilaterally rhytmical.
BACLOFEN AND CLONAZEPAM OVERDOSE IN A PATIENT WITH CHRONIC NECK AND SHOULDER PAIN

Delta and sparse sharp waves in the frontal regions. There were no signs of status epilepticus.

Conclusion
The present case demonstrates the importance of considering baclofen overdose in cases of severe coma in combination with generalized seizures, sinus bradycardia and normal blood pressure. Management of severe baclofen intoxication should be done at an ICU, where there is access to mechanical ventilation, forced diuresis, monitoring of vital signs with compensation of circulation, oxygen saturation and laboratory findings. Despite severe clinical manifestation of baclofen intoxication, the overall outcome is generally good.

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References