Acute hydrocephalus secondary to carbon monoxide poisoning

Hidrocefalia aguda secundaria a intoxicación por monóxido de carbono

Dear Editor:

Carbon monoxide (CO) poisoning is one of the most common causes of morbidity and mortality by poisoning. This type of toxicity results from tissue hypoxia and CO-mediated damage at the cellular level. Neurological sequelae constitute the principal cause of associated morbidity. The development of acute hydrocephalus is an extremely rare complication. We present a case of CO poisoning in an adult male who developed acute obstructive hydrocephalus secondary to bilateral cerebellar oedema.

The patient was a 38-year-old man with a medical history of hypothyroidism treated with thyroid hormone replacement therapy. He also presented personality disorder and had attempted suicide several times. He was found unconscious inside a car inhaling exhaust gases, and beside him were 2 empty blister packs of benzodiazepines. Upon arrival at the emergency department, the patient had a low level of consciousness (Glasgow Coma Scale = 3) and mid-dilated miopic pupils which were poorly reactive; as a result, he underwent orotracheal intubation. A cranial CT (Fig. 1) revealed diffuse hypodensities in both cerebellar hemispheres, basal temporal white matter bilaterally, both internal capsules, and the globus pallidus. The patient was admitted to the intensive care unit (ICU). A urine toxicology test revealed benzodiazepines and methadone, while CO-oximetry showed carboxyhaemoglobin levels of 23.6%. The patient was therefore mechanically ventilated with 100% oxygen until his carboxyhaemoglobin levels decreased to 0.9%, approximately 6 hours later. In the 48 hours after being admitted to the ICU, the patient’s neurological symptoms improved to the point where he was able to obey simple instructions, although he remained drowsy. However, 72 hours after admission to the ICU, the patient’s state of consciousness deteriorated suddenly. An additional cranial CT (Fig. 1) revealed severe hydrocephalus affecting the lateral ventricles and third ventricle, resulting in a significant mass effect. The patient underwent emergency surgery: first, an external ventricular drain was put in place, from which exuded a clear liquid under high pressure, and then, a decompressive craniectomy of the posterior fossa was performed to relieve cerebellar herniation. After surgery, the patient remained in coma for the next 5 days (flexion—extension of both upper limbs was the only response to painful stimuli). A follow-up cranial MRI scan performed 4 days after surgery (Fig. 2) showed multiple cerebral infaracts in an early subacute stage, extensively affecting the limbic system, hippocampus, fornix, and basal temporal area bilaterally. Patchy areas of small bilateral cortical infarcts could also be observed in the frontal and parietal lobes, as well as extensive infarcts in both cerebellar hemispheres, with the most damage occurring in the territory of the superior and anterior inferior cerebellar arteries; no signs of hydrocephalus were seen. Taken as a whole, these findings suggest multiple anoxic-ischaemic encephalic lesions secondary to CO poisoning.

As of the fifth day after surgery, our patient’s neurological symptoms had improved progressively, reaching an adequate level of consciousness and showing no focal neurological signs. Three days later, hydrocephalus resolved and the external ventricular drain was removed after remaining closed for 48 hours without visible neurological deterioration.

The clinical presentation and radiological findings of CO poisoning vary greatly. There are 3 mechanisms by which CO damages the central nervous system. Firstly, CO directly causes diffuse hypoxic-ischaemic encephalopathy, which predominantly affects grey matter. Second, although to a lesser extent, it may also cause focal lesions to the cerebral cortex, especially the hippocampus and temporal lobes. Cortical damage may manifest as transient vasogenic oedema or as a necrosis with infarct areas in the absence of cerebral arterial occlusion. Third, it may cause white matter demyelination. This finding, normally undetectable in the acute phase of intoxication, is considered the cause of late-onset neuropsychiatric syndrome. This syndrome generally develops after a lucid interval, and the most common symptoms are mental deterioration (amnesia, cognitive dysfunction), emotional disorders (depression, anxiety, mutism), urinary and faecal incontinence, and motor disorders (gait alterations, parkinsonian symptoms). The globus pallidus is the structure most frequently affected by CO poisoning; damage is usually immediate, bilateral, and predominantly affects the anterior 2 thirds of this structure. Occasionally the rest of the basal ganglia are affected (putamen, caudate nucleus, thalamus); in this case, the lesions are typically asymmetrical. The brainstem and the cerebellum are less...
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Figure 1  (A) Diffuse hypodensity in both cerebellar hemispheres, basal temporal white matter bilaterally, both internal capsules, and the globus pallidus. (B) Severe dilation of the lateral ventricles and third ventricle (the fourth ventricle was normal), which results in severe mass effect leading to effacement of the convexity sulci and a decrease in the size of the basal cisterns.

Figure 2  Multiple early subacute cerebral lesions which extensively affect the limbic system, as well as extensive infarcts throughout the cerebellar hemispheres.

frequently affected; thus, damage to these areas reflects more severe poisoning as these posterior structures are more resistant to hypoxia.1

Acute hydrocephalus is an extremely infrequent complication of CO poisoning which has previously been described in only 2 paediatric patients2,3 and one adult.4 In every case, the patients experienced an initial clinical improvement followed by rapid neurological deterioration after 24 to 72 hours. All neuroimaging tests revealed obstructive hydrocephalus secondary to bilateral cerebellar oedema with compression of the fourth ventricle. In the case of the adult patient, hydrocephalus resolved spontaneously and was accompanied by the recovery of an adequate level of consciousness within the next 72 hours, while in the paediatric patients progression was slow, leading to death in one of them, despite aggressive surgical treatment with ventriculostomy. These cases along with the current one suggest that, in addition to the structures most typically injured, there are others at the level of the brainstem and cerebellum which are highly susceptible to oedema caused by the cytotoxic effects of CO.

In conclusion, although the mechanism of the development of acute obstructive hydrocephalus is well known, this case is unusual by reason of being secondary to CO poisoning. Therefore, hydrocephalus should be suspected when acute neurological deterioration is observed in the days following CO exposure.

References

Sexsomnia and sleep eating secondary to sodium oxybate consumption

Dear Editor:

In a recent article, Ariño et al. reported 4 cases of sexsomnia, a parasomnia associated with confusional arousals during non-REM (NREM) sleep. Sexsomnia (sleep-related abnormal sexual behaviours) has been included in the most recent version of the International Classification of Sleep Disorders (ICSD-3) as a subtype of parasomnia.

Some of the actions considered to fall under this category are violent or prolonged masturbation, sexual harassment and/or assault (of minors or adults), initiation of sex (irrespective of the menstrual status of the bed partner), and loud vocalisations, often sexual in nature, while sleeping. The fundamental characteristic of this disorder is amnesia in regard to the episode the next morning or if the patient wakes up during the event. Episodes tend to occur in or near bed or wherever the patient sleeps; sexsomnia is rare among patients with sleepwalking who leave the bed or bedroom.

Such parasomnias as somnambulism, sleep-related eating disorder, and catathrenia have been described as side effects of sodium oxybate (SO) in the treatment of narcolepsy.

We report the case of a patient who presented sexsomnia and sleep-related eating disorder secondary to treatment with SO, an association which has not been described in the literature to date.

The patient was a 41-year-old woman with a diagnosis of narcolepsy with mild cataplexy. We initiated treatment with modafinil in increasing doses up to 300 mg/day; as the patient showed no response, she was treated with SO (4.5 g divided into 2 nightly doses), adding modafinil on request (100-150 mg). The reaction to this therapy was positive and our patient has since been able to lead a completely normal life: she started to work as a substitute teacher while studying for state qualifying exams.

Two years after treatment, the patient had an episode of self-limiting confusional arousal as a result of taking 4.5 g of SO in a single dose. However, treatment was not suspended.

A few months later, the patient reported sleep-related sexual episodes. At that point she had been taking 4.5 g of SO nightly for 1 year and 11 months. After taking the second dose of 2.25 g of SO, between 2.5 and 4 hours after the first, she sexually assaulted her bed partner (her husband), although she remembered nothing about it the next day. Her husband told her that she behaved strangely and aggressively; this behaviour was mostly accompanied by unintelligible sleep talking, but when her speech could be understood it was largely obscene. The sex act was only occasionally consummated; other times, after the sexual assault, the patient jumped out of bed and went to the kitchen, opened the refrigerator, and voraciously ate whatever she found there, although she showed a preference for carbohydrates.

She consumed no alcohol, drugs, or other medication either before or during the period during which these episodes occurred.

The sexsomnia and sleep-related eating disorder lasted for 3 weeks, after which SO treatment was temporarily discontinued resulting in immediate cessation of symptoms. SO treatment was slowly reintroduced and symptoms did not reappear.

Confusional arousal disorders during NREM sleep are often accompanied by disinhibition of such primitive behaviours as eating, having sex, or aggressiveness. Tassnari et al. proposed the hypothesis of a central pattern generator. These pre-established patterns of behaviour (whether eating, sexual, or defensive behaviour) are normally inhibited by the prefrontal cortex during wakefulness. Confusional arousal disorders are thought to be caused by a number of factors including a dissociation between different brain regions, activation of the central pattern generator, sleep inertia, and sleep-state instability.

In our view, SO may have the side effect of confusional arousals (including sexsomnia) probably due to the increase in slow wave sleep (stage 3 of NREM sleep).

References


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