Delayed encephalopathy after acute carbon monoxide poisoning

Akut karbonmonoksit zehirlenmesi sonrası gecikmiş ensefalopati

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ABSTRACT
Carbon monoxide poisoning is a major cause of death following attempted suicide and accidental exposures. Although clinical presentation depends on the duration and the intensity of exposure, the assessment of the severity of intoxication is difficult. A small percentage of patients who show complete initial recovery may develop delayed neurological deficits. Delayed encephalopathy after acute carbon monoxide poisoning is a rare and poor prognosis neurologic disorder and there is no specific treatment. We present a case with early onset of delayed encephalopathy after acute carbon monoxide poisoning with typical cranial imaging findings in a child with atypical history and clinical presentation.

Key words: Carbon monoxide, intoxication, encephalopathy

INTRODUCTION
Delayed encephalopathy after acute carbon monoxide poisoning (DEACMP) is a group of neuro-psychological disorders that occur days or weeks later after the disappearance of the symptoms of acute carbon monoxide (CO) poisoning [1,2]. In DEACMP the lucid interval before appearance of neurologic sequel varies from 2 to 40 days in adult patients. [1]. It has a poor prognosis and there is no a specific treatment [2]. Herein we present early onset of DEACMP with typical magnetic resonance imaging (MRI) findings following carbon monoxide poisoning in a child with atypical history and clinical presentation.

CASE REPORT
A 3.5 year-old boy was found unconscious on the tandoor, which is a traditional Turkish coal arger. He was admitted with second-degree burns on upper and lower extremities. On further enquiry, it was revealed that he had fallen over the top of a tandoor and nobody knew for how long he remained there. So the probability of CO poisoning was suspected. Biochemical tests including kidney function tests (urea, creatinine, sodium, potassium) and liver function tests were within normal limits. He was managed conservatively regarding CO poisoning and treated for his burns, in plastic surgery clinic. On the fourth day he had focal seizure and fell unconscious. The patient was transferred to pediatric intensive care unit. On neurological examination, his light reflex was nonreactive and he had spastic quadriparesis. Cranial MRI examination on the same day, revealed peculiar bilateral, symmetrical T2 hyper intensity in the basal ganglia, thalamus and subcortical white matter of the occipital lobes.

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Seizures stopped after administration of antiepileptic therapy.

Since hyperbaric-oxygen therapy is unavailable in our center, he was followed with nasal oxygen support during his comatose state, which in turn made no change in his neurological status. Currently he is being followed by physical therapy program.

**Figure 1.** T2 weighted MRI image obtained four days after acute CO intoxication shows bilateral, symmetrical hyperintensity in the basal ganglia, thalamus, and subcortical white matter of the occipital lobes

**DISCUSSION**

Prompt diagnosis of CO poisoning requires strong clinical suspicion and detailed patient history indicating duration of the exposure. There are many reasons that can cause CO poisoning. Delayed encephalopathy after acute carbon monoxide poisoning has a poor prognosis. In adult patients the lucid interval before appearance of neurologic sequelae varies from 2 to 40 days in DEACMP [3]. Kim et al reported that neurological symptoms which are preceded by a lucid interval of two to three weeks after the recovery of the acute stage may be observed in delayed encephalopathy of carbon monoxide poisoning [4]. In present case, neurological signs appeared on the fourth day and the patient was transferred to the intensive care unit immediately.

The MRI neuroimaging findings in CO poisoning may be detected before appearance of clinical symptoms. MRI is a highly sensitive method to demonstrate the neuroimaging abnormalities after CO poisoning. Changes may visible as early as 1 hour post-exposure. Acute CO poisoning typically shows deep gray matter involvement, most commonly in the globus pallidii, putamen and thalami, occasionally with a hemorrhagic component [3]. In present case, MRI examination was performed as the neurological signs first appeared, and bilateral, symmetrical T2 hyper intensity in the basal ganglia, thalamus and subcortical white matter of the occipital lobes was observed.

Herein we would like to specify that CO poisoning should also be keep in mind in patients with burns. To our knowledge, this case is one of the pediatric case in the literature, presenting with the earliest clinical and neuroimaging findings of DEACMP.

**REFERENCES**