Methanol-induced parkinsonism and cerebral vasculopathy due to perfume inhalation

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ABSTRACT

Background Methanol (methyl alcohol) is a form of toxic alcohol that is found in illicit alcohol as well as household products such as solvents and paint removers. The most common cause of methanol poisoning is through ingestion of adulterated alcohol; however, other routes of poisoning may also occur including cutaneous exposure and, rarely, inhalation.

Methods/results We are reporting a case of a young woman with vision loss, parkinsonism and widespread cerebral artery spasms due to methanol inhalation from domestically made perfume.

Conclusion Our case highlights the increased need for awareness on the part of the public and health authorities with regard to the manufacturing and use of homemade perfumes produced with poorly processed alcohol having a high methyl alcohol content.

INTRODUCTION

Methanol is an alternative to ethanol because it is cheaper and easier to manufacture. Its illegal oral consumption is the most commonly reported route of toxicity in the literature. Methanol poisoning is thought to cause severe morbidities such as vision loss, metabolic alterations and irreversible neurological dysfunction, and it may even lead to mortality. We are reporting an unusual presentation of incidental methanol intoxication through the inhalation of homemade perfume during the manufacturing process.

CASE REPORT

A woman in her 30s living in a rural area of Saudi Arabia was referred to our tertiary care facility for the management of sudden vision loss and parkinsonism. The patient makes a living by manufacturing local perfumes at home, and to compensate for increased demand for her product, she admitted to using adulterated alcohol from a different local vendor. One day after producing a large batch of perfume, she experienced sudden vision loss and became encephalopathic. Subsequently, she was admitted to her local hospital as a case of encephalopathy with optic neuritis, and she required intubation. Due to the lack of the aforementioned noted history, she was misdiagnosed as having autoimmune encephalitis, for which she received a course of pulse methylprednisolone followed by intravenous immunoglobulin G (IVIG). Her initial laboratory investigations (including cerebrospinal fluid analysis) were unremarkable, and the brain MRI showed bilateral and symmetrical areas of bright T2 signals involving the basal ganglia with surrounding oedema. The lentiform nuclei demonstrated blooming in T2-weighted images donating blood degeneration products, likely secondary to necrosis. The frontal and occipital lobes showed bilateral, symmetrical, subcortical, hyperintense T2 signals with heterogeneous contrast enhancements in addition to enhancement of the optic nerves bilaterally (figure 1).

On discharge from her local hospital, the patient showed improvement only in her level of consciousness with no significant change.
in her visual acuity. She became apathetic and bradykinetic, which led to her referral to our tertiary centre.

The patient was hypomemic, hypophonic and apathic with minimal verbal output. She had bilateral polyminimyoclonus in her hands. The higher mental functions and cranial nerves were intact, aside from bilateral optic nerve atrophy and visual acuity with appreciation of hand motions only. The power and deep tendon reflexes were normal. Significant rigidity and bradykinesia were noticed. A sensory examination was unremarkable to all modalities as well as cerebellar examination. Her gait revealed a decreased arm swing bilaterally with a shuffling gait and mild postural instability. The estimated unified Parkinson’s disease rating scale (UPDRS) III was 50.

Laboratory investigations in our hospital, including a haematological profile, renal profile and hepatic profile, were unremarkable. An antinuclear antibody screening, a pyruvate kinase test, lactate levels, and a Wilson workup were normal. A visual evoked potential test showed prolonged P100 bilaterally.

Repeated MRI brain scans showed evolution of the previously mentioned changes with the development of patchy cystic enhancement and microhemorrhages in the basal ganglia, as well as in the frontal and occipital lobes bilaterally. There was enhancement of the bilateral optic nerves and optic chiasm (figure 2). Brain magnetic resonance angiography demonstrated multiple instances of long segmented narrowing along the bilateral A2, A3 and right A1, M2 as well as mild stenosis of left M2. PET-CT of the brain FDG showing symmetrical FDG uptake within the caudate nucleus with marked symmetrical reduced FDG activity within the lentiform nucleus and mild hypometabolism seen in the occipital lobe bilaterally. PET-CT of the brain and F18-DOPA scan showing reduced tracer activity involving the caudate nuclei bilaterally as well as the inhomogeneous, asymmetrical activity involving the putamen and globus pallidus bilaterally more pronounced on the left. FDG, florodeoxyglucose; F-DOPA, 6-[18F]-L-fluoro-L-3, 4-dihydroxyphenylalanine; MR, magnetic resonance; PET, positron emission tomography.

Figure 2 (A) MR angiography of the brain demonstrate multiple long segmented moderate smooth narrowing along the bilateral A2, A3 and right A1, M2 as well as mild stenosis of left M2. (B) PET-CT of the brain FDG showing symmetrical FDG uptake within the caudate nucleus with marked symmetrical reduced FDG activity within the lentiform nucleus and mild hypometabolism seen in the occipital lobe bilaterally. (C) PET-CT of the brain and F18-DOPA scan showing reduced tracer activity involving the caudate nuclei bilaterally as well as the inhomogeneous, asymmetrical activity involving the putamen and globus pallidus bilaterally more pronounced on the left. FDG, florodeoxyglucose; F-DOPA, 6-[18F]-L-fluoro-L-3, 4-dihydroxyphenylalanine; MR, magnetic resonance; PET, positron emission tomography.
Symptomatic therapy consisting of carbidopa–levodopa and pramipexole was started with a gradual increase in the dosages.

During a follow-up visit a month later, the patient exhibited no change in her visual symptoms, but she reported an improvement in her activities of daily living which corresponded to her UPDRS improvement by 20%.

**DISCUSSION**

This report represents an unusual presentation of incidental methanol intoxication through inadvertent inhalation while manufacturing homemade perfume. It also highlights an additional, yet unrecognised finding of cerebral artery spasms as a squeal of methanol poisoning.

Methanol intoxication can cause severe visual disturbances, metabolic alterations, gastrointestinal complaints and irreversible neurological dysfunction, and it may even be fatal. Symptoms may manifest after a latency period of a few hours after exposure (as seen with our patient) or may develop days later. The damage is due to alcohol dehydrogenase’s oxidation of methanol, which causes its conversion to formaldehyde, leading to formic acid production—a toxic metabolite. Formic acid causes toxicity by alterations in the cytochrome c oxidase complex at the mitochondria’s terminal end of the respiratory chain. This change causes disruption of the mitochondria, which are represented heavily in the optic nerves, retina and putamen.

MRI brain findings in methanol toxicity include haemorrhagic putaminal necrosis, diffuse white matter lesions, cerebellar or brainstem lesions that may enhance, and intracerebral or intraventricular haemorrhaging. In a case series following 58 patients with methanol intoxication, 33 cases suffered from severe visual disturbances, and brain imaging revealed bilateral optic nerve enhancement, while 45 cases revealed bilateral putaminal ischaemic changes or necrosis. All of these cases experienced extrapyramidal symptoms. Of note, there was no correlation between the degree of optic atrophy and putaminal involvement. Other findings mentioned included asymmetric putaminal involvement, caudate involvement and subcortical white matter oedema.

What makes our case significant is the presence of multiple cerebral vessel stenosis, which has not been reported in the literature previously (figure 2A). The mechanism of methanol-induced vasospasm was studied in the late 1990s as a response to methanol exposure in canines. The study reported a similar finding to ethanol-induced vasospasm, hypothesised to be secondary to its direct action on the smooth muscles of the cerebral vasculature in opposition to alteration in metabolic demand. On autopsy, brain infarctions have been previously reported in humans in cases of methanol intoxication; however, vessel stenosis has not been highlighted.

This is the second case reported to have presynaptic dopaminergic denervation as manifested by reduced tracer activity in the F-DOPA scan (figure 2C). The first case was a female presenting with delayed-onset parkinsonism—dystonia and hypophonia after methanol intoxication, as reported by Franquet et al. In our patient, [18F] fluoro-L-DOPA PET imaging confirmed dopaminergic nigrostriatal neuron functional impairment via reduced tracer activity involving the basal ganglia, which explains the parkinsonian features induced by methanol toxicity. These findings were concurrent with the patient’s MRI findings that were reported previously (figure 2C).

Parkinsonian symptoms resulting from methanol poisoning have been managed by the use of dopaminergic agents, as was the case with our patient who showed a fair response to levodopa–carbidopa and pramipexole; however, the literature showed conflicting evidence in response to treatment.

Our case highlights the increased need for awareness on the part of the public and health authorities with regard to the manufacturing and use of homemade perfumes produced with poorly processed alcohol having a high methyl alcohol content.

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