Lesions of basal ganglia due to disulfiram neurotoxicity

D Laplane, N Attal, B Sauron, A de Billy, B Dubois

Abstract

Three cases of disulfiram induced Parkinsonism and frontal lobe-like syndrome associated with bilateral lesions of the lentiform nuclei on CT scan are reported. Symptoms developed either after an acute high dose of disulfiram (one case) or after several days to weeks of disulfiram treatment (two cases) and persisted over several years in two patients. These observations suggest that basal ganglia are one of the major targets of disulfiram neurotoxicity. The mechanisms of the lesions of basal ganglia may involve carbon disulfide toxicity.

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Various central neurological manifestations following disulfiram intoxication have been described.1–10 They include symptoms suggestive of basal ganglia dysfunction, such as parkinsonism and catatonia.4–10 Lidy et al reported hypodensities of basal ganglia on CT scan after acute disulfiram intoxication,9 but no clear relation has been established between parkinsonism or catatonia due to disulfiram toxicity and basal ganglia lesions. A recent extensive review of disulfiram induced catatonia failed to mention any possible implication of such lesions in the catatonie syndrome.5 We report three cases in which symptoms indicative of basal ganglia dysfunction were associated with documented lesions of the basal ganglia after disulfiram intoxication.

Case reports

Case 1

A 42 year old black woman from Guyana with a five year history of alcohol abuse had been treated on two occasions with disulfiram. On the evening of admission she had voluntarily taken 75 disulfiram 500 mg tablets, half a litre of champagne, 5–10 diazepam 10 mg tablets, and 40 B1–B6–B12 vitamin pills. Initially alert, she was admitted to the hospital emergency unit, where she refused gastric lavage. Results of neurological examination were described as normal. Within five days she progressively became somnolent and dysarthric, with limited comprehension for simple commands. She experienced nightmares and auditory hallucinations, along with paranoid delirium. These symptoms subsided over the following days. Results of electroencephalography were normal. Laboratory test results were within normal limits, except for moderate macrocytic anemia.

Three weeks after her admission a generalised tremor of the extremities was observed at rest, and she developed dysphagia and difficulties in swallowing consistent with pseudobulbar palsy. She was dystrophic, with a nasalised and monotonous voice and buccofacial apraxia. Imperious micturition was noted. The plantar reflexes were flexor and the cutaneous abdominal reflexes were not elicited. The facial reflexes were brisk. She kept her eyelids wide open, with rare blinking. Additional neurological findings included signs of polyneuropathy, with sural biopsy showing acute axonopathy. She received carbamazepine (200 mg daily) for painful dysesthesias, as well as vitamins. Results of tests for B12 vitamins, folates, blood ammonia, and urinary porphyrins and serum and urine screens for heavy metals were normal. Spinal fluid examined on day 9 was normal. An EEG showed diffuse symmetrical slowing in the theta and delta range, without focal or epileptiform abnormalities. On about day 32, her condition stabilised and then gradually improved. A few days later myoclonic jerks affecting both arms were noted; these subsidised in the following week.

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Figure 1 Case 1: horizontal CT scan showing small bilateral lucencies in the putamen.
CT scans performed one and two months after admission revealed almost symmetrical bilateral low density lesions of the basal ganglia, occupying the putaminal nuclei (fig 1). On discharge two months after admission the patient had totally recovered.

Case 2
A 23 year old man had been an alcoholic for eight years; he had been started on disulfiram 1 g daily eight weeks before admission and had abstained from alcohol. His personality was considered to be sociopathic. His history included several admissions to hospital for alcoholic treatment and mild depression.

Seven weeks after disulfiram had been started he was noted to have a progressively worsening staggering gait. He was thought to have resumed taking alcohol, and disulfiram was increased to 2 g daily. In 48 hours he was admitted to an intensive care unit unresponsive. Examination did not reveal any focal neurological signs. The plantar reflexes were flexor and the deep tendon reflexes were absent. The patient was put on a ventilator because of cyanosis and bronchial obstruction. A chest radiograph did not show any pulmonary infection. Arterial blood gas levels (room air) revealed a \( \text{PaCO}_2 \) of 58 mm Hg, \( \text{PaO}_2 \) 31 mm Hg, and \( \text{pH} \) 7.27. Serum alcohol was negative. Laboratory results showed leucocytosis and diabetic ketoacidosis. Treatment with intravenous insulin was started. A first lumbar puncture was traumatic, with protein 0-3 g/100 ml. An EEG showed diffuse slowing, without any paroxysmal activity. Disulfiram was withheld.

On day 3 the patient was alert and extubated. Insulin dosage was reduced. The next morning, ventilation was again required because of increasing impairment of consciousness. Facial myoclonus was noted, and a second EEG showed generalised epileptiform waves. Phenobarbital 20 mg daily and nitrazepam 3 mg daily were started. A CT scan performed on day 5 showed bilateral low density areas occupying the lentiform nuclei and extending onto the internal capsules. On day 10 a chest radiograph showed left lobar atelectasis and pleural effusion at the base of the lung. Pleural puncture showed bacterial infection with *Klebsiella* sp, *Staphylococcus aureus* and *Escherichia coli*, for which treatment with penicillin, metronidazole, and tobramycin was begun. Tracheotomy was performed.

Within three weeks the patient’s vigilance, chest, and physical conditions gradually improved, as did EEG and laboratory findings. Blood glucose concentration had normalised and insulin therapy was withheld. Successive lumbar punctures were normal.

One month after admission the patient became agitated and emotionally labile. Meprobamate (800 mg day) was given. The facial reflexes were brisk and the velar reflex was absent. No plantar reflex was elicited. Additional findings disclosed signs consistent with peripheral neuropathy, with sural biopsy showing diffuse axonopathy. Laboratory findings showed normal concentrations of serum B12, folate, and urinary porphyrins. Serologic studies for syphilis, HBs antigen, antinuclear bodies and serum complement were normal. CT scans performed one and two months after admission showed no change (fig 2). On his discharge two months after admission, the patient’s condition had notably improved, but mild proximal extrapyramidal rigidity was noted.

Seven months later, on follow up, the patient had not resumed alcohol consumption and was not taking disulfiram. Examination showed signs of parkinsonism, with extrapyramidal rigidity of both arms, facial akinesia, a typical abnormal posture, and a slowing of the gait. There was no tremor. The patient was severely dystarthric, with tonic stuttering but no palilalia. Blepharospasm and dystonic posture of the left hand were noted. The facial reflexes were brisk and the plantar reflexes were absent. Grasping and sucking reflexes were not elicited. The tendon reflexes were brisk in the arms and absent in the legs. Additional findings showed residual signs of peripheral neuropathy. The CT scan showed no change and cerebrospinal fluid was normal. No treatment was given for parkinsonism. One year, and seven months later neurological findings were comparable except for a central reflex neurogenic bladder, for which he received treatment.

Case 3
A 52 year old man with a history of ethanol abuse had received disulfiram (500 mg daily for several months) 12 years previously, at which time he stopped consuming alcohol. He was being treated with phenobarbital for epilepsy. His history did not reveal any psychiatric disease.

He was admitted to hospital because of faintness. He was soon noticed to have strange behaviour, which had been observed by his
Pallidal areas on a CT horizontal slice. Lesions and right hemisphere performed that lesions blunted affect. Scales (only achieved) were Luria’s Minnesota Personality inventory. Wechsler calculation, was normal. There was no impairment of learning and acquisition (Wechsler MS MQ was 100) either of old or recent personal memory. Mental control subtests were well performed. Verbal fluency was very poor for the enumeration of words beginning with “m” (one within 60 seconds). The elaboration of new strategies and the sequential programming of activities were disturbed, as shown by the Wisconsin card sorting test (only two categories achieved) and Luria’s graphic series. Personality (evaluated by the Minnesota multiphasic personality inventory) was not impaired; depressive rating scales were negative, but he showed blunted affect.

CT scans showed bilateral low density lesions that projected onto the pallidal and putaminal areas (fig 3). An MRI scan was performed five months later and showed a right side lesion involving parts of the putamen.

Discussion
In the three cases that we report, symptoms suggestive of basal ganglia disease were observed after disulfiram treatment. Two patients developed parkinsonism and pseudobulbar-like syndrome and one developed blepharospasm. The third patient did not show motor disorders but exhibited “loss of drive” or more specifically “loss of psychic self activation.” This behavioural disorder, which resembles the negative symptoms of schizophrenia or the mild forms of catatonia, has been reported recently to result from basal ganglia lesions.

The symptoms exhibited by the patients were clearly related to disulfiram and to no other pathological condition in patients 1 and 3. Patient 1 developed parkinsonian-like symptoms after taking disulfiram together with alcohol. Acute alcohol intoxication or withdrawal may provide parkinsonism in chronic alcoholics with underlying parkinsonian pathology, but lesions of the lentiform nuclei on CT scan due to alcohol alone have not been described. Although patient 3 was only seen 12 years after disulfiram treatment, it was clear from his relatives that his behavioural disorders started at that time but were not considered severe enough to need medical care. The relation between disulfiram and cerebral insult may be more problematic in patient 2, who suffered from hypoxia due to pulmonary infection, since the vulnerability of basal ganglia lesions to anoxic injury has long been recognised. However, hypoxia in this patient was moderate, as shown by blood gases, and probably insufficient to induce signs of brain hypoxia. In addition, bilateral ganglionic low density lesions were seen on CT scan only five days after the patient was intubated, and these lesions usually take longer to develop after anoxia. Indeed, in recently reported cases, CT scan performed a few days after cerebral anoxia showed high density ganglia lesions.

CT scans showed bilateral lesions of basal ganglia in all patients. The lesions were confined to the lentiform nuclei, particularly affecting the putamen. In patient 3 who underwent CT scan long after disulfiram treatment the lesions remained stable over several years, which is evidence in favour of an irreversible mechanism. Interestingly, the lesions were similar in all three cases, whatever the dosages and duration of disulfiram, which ranged from 500 mg daily to an acute intoxication with 37.5 g. This is consistent with previous observations, showing that the incidence of disulfiram intoxication does not differ between groups receiving various dosages. As disulfiram has a very long half life these phenomena may relate to accumulation of the drug because impaired hepatic function in patients with liver damage induced by alcohol.

Disulfiram induced parkinsonism and catatonia are well documented. In contrast, basal ganglia lesions due to disulfiram have not been commonly reported. Our data suggest that disulfiram intoxication may be related to such lesions, as is the case with post ence-
phalitic parkinsonism, carbon monoxide intoxication, or cerebral anoxia, the negative CT scans in recently reported cases of catatonia or parkinsonism due to disulfiram may be because these were less severe cases. The mechanisms underlying disulfiram induced parkinsonism or catatonia are still controversial. One of the main hypotheses is that the lesions result from the toxicity of carbon disulfide, one of the disulfiram metabolites. The symptoms induced by carbon disulfide, which include parkinsonism and peripheral neuropathies, resemble those of disulfiram intoxication. Patients 1 and 2 developed peripheral neuropathy after receiving disulfiram, and morphological evidence has been provided for carbon disulfide toxicity in disulfiram neuropathy. Moreover, histopathological lesions of the globus pallidus and substantia nigra have been observed after chronic exposure to carbon disulfide in animals.

Other authors have suggested involvement of brain dopaminergic transmission in disulfiram neurotoxicity, and in mice the toxic effects of MPTP on nigrostriatal dopaminergic systems was enhanced by pretreatment with disulfiram. Indeed, disulfiram induces enzymatic blockade, which impairs the ability to eliminate free radicals. Interestingly, one of the main hypotheses about the causative factors of Parkinson’s disease is that the impairment of dopaminergic nigrostriatal systems results from production of free radicals. In the experiment on mice, however, disulfiram treatment without MPTP did not impair the dopaminergic systems significantly. Nevertheless, it is difficult from the present data totally to exclude a dopaminergic mechanism, since we have no information on the functioning of the substantia nigra in the patients. A similar case has recently been reported.

In conclusion, our case reports provide evidence that disulfiram induced extrapyramidal and behavioural disorders may be related to lesions of basal ganglia. It would thus be interesting to perform MRI scans in patients with disulfiram induced neurotoxic reactions. Our observations are consistent with the hypothesis that disulfiram neurotoxicity is mediated through carbon disulfide. They emphasise that basal ganglia lesions are a major site of action for disulfiram neurotoxicity. As indicated from previous experiments and in agreement with initial case reports, these data suggest that a separate disruption of parallel basal ganglia-thalamocortical circuits may be involved in the behavioural and motor disorders shown by patients with basal ganglia lesions.

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Bourneville's tuberous sclerosis: “When the cat is away...”

Whilst deputising for Mr L J F Delaisseau at the Salpêtrière on 18 July 1867, Désiré Magloire Bourneville (1840–1909) observed a child with the syndrome which now bears his name. He published the case promptly⁠¹ and thereby his name gained eponymous precedence over Dr Hartdegen who made similar observations at the same time.² Bourneville’s patient was Marie aged three, born of parents without neurological abnormality or consanguinity.

“... breast fed by a wet nurse until 14 months. During this time she may have had several convulsions restricted to the eyes. At two years seizures appeared, during which the arms shook and turned slightly. It was particularly in the head. Marie never walked or talked. She gradually deteriorated... a hopeless case.

The eyes are dull... Rosaceous and pustular acne(sic) of the face; a confluent vesiculopapular rash over the nose, cheeks and forehead; numerous small mollusca on the nape of the neck, which is abnormally short... The right arm is paralysed, not complete... The left leg is longer and heavier than the right. The right thigh is adducted and flexed on the pelvic; the foot is flat, in varus, and violaceous... The legs are bent and crossed in bed. Constant dribbling... The seizures came in series... April 20th–30th fits... enema of bromide of camphor 2 grains.”

After further fits the child died on 7 May at 3 am. At necropsy the brain weighed 1000 gm. He saw lesions: “rounded islets, forming protuberances of vari-
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