Accidental ethyl mercury poisoning with nervous system, skeletal muscle, and myocardium injury

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SUMMARY Four case reports are presented of patients who ate the meat of a hog inadvertently fed seed treated with fungicides containing ethyl mercury chloride. The clinical, electrophysiological, and toxicological, and in two of the patients the pathological data, showed that this organic mercury compound has a very high toxicity not only for the brain, but also for the spinal motoneurones, peripheral nerves, skeletal muscles, and myocardium.

Use of organic mercury compounds as fungicides has been noted in several countries to produce severe poisoning, occasionally epidemic in character. We had the opportunity to examine the first cases of such poisoning ever to have appeared in Romania. They showed some features not yet described in the literature on poisoning with various organic mercury compounds.

Case reports

In August 1974, a mother with her three children was referred to our clinic by a provincial hospital where they had treated for 17 days.

CASE 1
This boy (MG), aged 15 years, one week before his first admission developed pain in the muscles and gait disturbance. He could neither stand nor walk. The tendon jerks were bilaterally hyperactive and there was a positive Babinski's sign on the left side, clonus of the left foot, and twitching of the thigh muscles. Severe cerebellar signs were noted—that is, ataxia and dysarthria. Within a week he developed dysphagia and aphonia, Babinski's sign appeared bilaterally, and the overall status deteriorated. On referral to this clinic we noted generalised muscle wasting, spasticity of the upper limb flexors and lower limb extensors, bilateral mydriasis, and impaired swallowing. The patient was drowsy, unresponsive to commands, and had loss of sphincter control. The clinical picture deteriorated even further: he developed a horizontal nystagmus, was agitated, and became comatose four days later. He required ventilatory assistance and died three days later from cardiac arrest, within one month of the onset of the illness.

The CSF was clear with 0.4 lymphocytes per mm³ and 0.28 g/l albumin. Haemoglobin was 10.88 g/dl. The leucocyte count was 13,500/mm³ with neutrophilia. The blood glucose was 8.6 mmol/l, urea 17.3 mmol/l, creatinine 248.5 μmol/l, sedimentation rate 55 mm in the first hour, 90 mm in the second hour. There were traces of albumin in the urine.

An electroencephalogram showed slow diffuse dysrhythmia with predominance of delta waves. General necropsy revealed foci of bronchopneumonia and an abscess of the left lung. The gross appearance of the brain was normal. All over the cerebral cortex but mostly in the calcarine cortex, there was nerve cell loss and a diffuse proliferation of neuroglia on microscopic examination of the brain.

In the midbrain and bulbar (especially the lateral nucleus) reticular formation, neuroglia activation with nerve cell satellitosis was apparent, extending to neuronal loss and formation of neuronophagic nodules.

Myelin preparations showed demyelination of the nerve fibres in the ninth and tenth cranial nerve roots (fig 1) In the cerebellum there was...
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Fig 1  Demyelination of nerve fibre bundles in the vagus nerve roots (case 1). Spielmeyer staining, original magnification ×100.

Cell wasting of the granular layer. Purkinje's cells were more spared, though in certain areas silver impregnation for neurofibrils showed empty basket-cells and torpedo-shaped Purkinje cell axons (fig 2). The lesions were evenly distributed over the neo- and paleocerebellar cortex. Throughout its length the spinal cord showed a highly abnormal picture of the ventral horns, with motoneurone loss and presence of neurophagocytic nodules (fig 3). Among the spared neurones many showed chromatolysis or were shrunken or hyperchromatic. The spinal and peripheral nerve roots were normal in appearance.

Fig 2  Empty basket-cells and torpedo-like axon of Purkinje cell in cerebellar cortex (case 1). Silver impregnation Bielschowsky, original magnification ×240.

The skeletal (intercostal) muscles showed zones of waxy segmental degeneration of the muscle fibre, with interstitial proliferation of fibroblasts (fig 4).

Fig 4  Intercostal muscle (case 1). Waxy degeneration of muscle fibre segments with interstitial fibroblast proliferation. Haematoxyline and eosine staining, original magnification ×100.

On microscopic examination of the other viscera, the kidney showed stasis and interstitial infiltration, with rarefaction of the glomerular elements here and there. The myocardium showed zones of interstitial chronic myocarditis with proliferation of fibroblasts but absence of sclerosis (fig 5). In the lung bronchopneumonic foci were found. The liver was normal.

CASE 2

This 10 year old boy (MM) had, five days before admission, developed walking difficulty, muscle pains, vomiting, and dysarthria. Neurological examination showed walking disability, weakness of the limbs, general hypotonia, horizontal nystagmus, deglutition impairment, severe ataxia, and increased tendon reflexes.
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Over the next two weeks he developed psychomotor agitation and a bilateral Babinski sign. He became unable to speak and entered a subcoma state.

He was still comatose when referred here. He could not swallow, the pupils were bilaterally dilated, and there was generalised muscle atrophy and loss of sphincter control. His condition deteriorated gradually. Within 10 days he developed decortication posturing, fever, and vomiting. He died from cardiac arrest within a month and a half of the onset of the disease.

The CSF was clear, with 0.8 lymphocytes/mm³; albumin 0.16 g/l; blood contained 10.22 g/dl haemoglobin. The white cell count was 10,000/mm³ with neutrophilia. Blood urea was at first 6.7 mmol/l and then increased to 19.7 mmol/l. Sedimentation rate was 27 mm/hr; urine albumin level was 3.3 g/l.

Electrocardiography showed S-T segment depression and T wave inversion in precordial leads. The electroencephalogram was of average voltage with lack of alpha rhythm and diffuse slow activity.

On gross pathological examination, the brain appeared normal. Microscopic examination of the cerebral cortex showed lesions which were identical to those found in case 1 but with a much more intense neuroglial proliferation that had resulted in formation of neuroglial nodules in the calcarine area (fig 6).

The caudate nucleus and the putamen showed an almost complete loss of the small cells with preservation of most of the large cells and diffuse proliferation of glia. In the brainstem there was glial proliferation in the midbrain periaqueductal grey matter. In the cerebellum cell loss was noted in the granular layer. The spinal cord showed alterations identical to those found in case 1.

Pathological examination of the viscera revealed acute inflammation of the renal pelvis, stasis and tiny lesions of interstitial nephritis.

Chronic interstitial myocarditis lesions were found as in case 1 (fig 7).

In the lung there was a picture of oedematous alveolitis. The liver was normal.

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**Fig 5** Interstital chronic myocarditis with fibroblast proliferation (case 1). Haematoxyline and eosine staining original magnification × 240.

**Fig 6** Calcarine cortex: disappearance of neurones with intense neuroglial proliferation and formation of neuroglial nodules (case 2). Nissl staining, original magnification × 100.

**Fig 7** Interstitial chronic myocarditis (case 2). Haematoxyline and eosine staining, original magnification × 240.

**CASE 3**

The mother (MT), a 48 year old farmer, when admitted to the hospital accompanying her two sons had no pathological sign whatever. Within 48 hours she developed gait disturbances with
progressive weakness in the lower limbs and muscular pains. Referred to this clinic, she was noticed to have spastic paraparesis, with bilateral Babinski sign and distal superficial hypoesthesia of the limbs. She was drowsy. Nine days later she developed headache and vomiting, her sight deteriorated and there was a horizontal nystagmus and intention tremor. She became agitated, confused and delirious. Within one month of the onset of symptoms she improved and was able to walk again. The pyramidal tract signs remitted and so did the mental disturbance. She continued to have headache and fatigue. The visual field was concentrically narrowed.

Routine laboratory tests failed to show any abnormality, except a slightly raised blood urea (7.3 mmol/l), while in the CSF there was a slight pleocytosis (15 lymphocytes/mm³) with an albumin level of 0.3 g/l.

Electrocardiography showed flattening of the T wave in precordial leads. An audiogram revealed slight bilateral perceptive deafness. The EEG tracing was of slightly decreased amplitude, with the alpha rhythm present and predominance of fast waves.

Two months later she was discharged home much improved. She was readmitted for a review four months later and was found to show only a concentric narrowing of the visual fields.

CASE 4

This 15 year old girl (MA), was admitted one week after her brother. Five days previously she had developed gait and speech impairment and a state of drowsiness. Her gait was spastic and slightly ataxic. Babinski’s sign was present on both sides. Speech was dysarthric.

When examined here she was found to have marked amblyopia with bilateral mydriasis and slow pupillary reflexes. She could walk only if supported. Her gait was spastic and she had spastic quadriaparesis, drowsiness and disturbed sphincter control. Three days later, vomiting, intention tremor, staccato speech, and slowness of cerebration developed. Within 10 days of admission the motor deficiency remitted, she regained the ability to distinguish the shape of things, though she still had concentric narrowing of the visual fields. She recovered progressively; her sight improved as she was now able to see at 5 m distance. She started to walk again and the cerebellar signs diminished.

The CSF contained no cells, albumin 0.16 g/l. Urine albumin was noted at a concentration of 0.5 g/l.

Her electrocardiogram showed sinus rhythm at 100 beats/min. In precordial leads the S-T segment was below the isoelectric line and the T wave was negative. There was evidence of left ventricular hypertrophy. The electroencephalogram showed slow dysrhythmia.

The diagnosis established during hospitalisation was organic mercury poisoning, and treatment with penicillamine 1 g/day for 21 days was started one month after admission. She was discharged greatly improved after a stay of two months in the clinic. When reviewed four months later she still had a mild intention tremor and concentric narrowing of the visual fields.

The clinical diagnosis of organic mercury poisoning was confirmed by the toxicological determinations performed on the blood, hair and urine immediately after admission (tables 1, 2).

Table 1  Mercury content of blood and hair

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
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<tbody>
<tr>
<td>Mercury level in whole blood (µg/ml)</td>
<td></td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>(maximum permitted 0.02 µg/ml)</td>
<td>0</td>
<td>5</td>
<td>1</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Mercury level in hair (µg/g)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(maximum permitted 6 µg/g)</td>
<td>__</td>
<td>542</td>
<td>160</td>
<td>152</td>
<td></td>
</tr>
</tbody>
</table>

Table 2  Mercury in the urine (µg/24 hr; maximum permitted 100-120 µg/24 hr)

<table>
<thead>
<tr>
<th></th>
<th>Period</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
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<tr>
<td>Mercury level in urine</td>
<td>Admission</td>
<td>57</td>
<td>126</td>
<td>660</td>
<td>165</td>
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<tr>
<td></td>
<td>10 days later</td>
<td>180</td>
<td>452</td>
<td>224</td>
<td></td>
</tr>
<tr>
<td>Mercury level after starting penicillamine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>After 7 days treatment</td>
<td>__</td>
<td>__</td>
<td>303</td>
<td></td>
</tr>
<tr>
<td></td>
<td>After 14 days treatment</td>
<td>__</td>
<td>__</td>
<td>193</td>
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</table>

Subsequent epidemiological data showed that 10 days or so before becoming ill the family had eaten the meat of a hog that over the past few days had shown signs of disease, reeling and falling. Some poultry died with the same signs. They had inadvertently been fed seed treated with fungicides containing alkyl mercury compounds —Cryptodin (a powder containing 2.5% ethyl mercury chloride) and F.B. (a powder containing 1% ethyl mercury chloride).

ELECTROPHYSIOLOGICAL INVESTIGATION

In the two surviving patients the following additional investigations were performed.
In cases 3 and 4 EMG recordings were obtained from the tibialis anterior, extensor digitorum brevis of the foot and the first dorsal interosseous muscle of the hands. They were all normal. Motor conduction velocity in cases 3 and 4 was measured in the median and ulnar nerves (segment elbow to wrist) and in the peroneal nerve (segment knee to ankle). Standard techniques were used with recording of the evoked responses in the abductor pollicis brevis, hypothenar, and extensor digitorum brevis muscles respectively.

Sensory conduction velocity was measured in the median and ulnar nerves. The stimuli were delivered to digits 1 and 5 respectively through ring electrodes. Recordings were made from the nerves at elbow and wrist. The results are shown in table 3.

The levels of mercury in the hair, urine, and (in lethal cases) in the nervous system (table 4) and viscera (table 5), were measured by a photometric method with dithizone after decay of the organic substances in an acid oxidant (Nestorescu's method). Unfortunately we were not able to use more reliable methods (atomic absorption and gas chromatography).

**Discussion**

In all our patients the symptoms appeared at least 10 days after ingestion of mercury contaminated foodstuffs, as reported also by other authors in organic mercury poisoning.2-5

The initial symptoms were headache, fever, diarrhoea, vomiting and myalgia, recalling especially the symptomatology in a report on poisoning by an ethyl mercury compound (ethyl mercury p-toluene sulphonanilide) in the 1961 Iraq epidemic.6

Although the neurological symptoms of poisoning by alkyl mercury compounds are well known especially after the 1953 Minamata epidemic, it might be of interest to attempt to correlate certain phenomena noticed in our patients with the anatomical lesions and the electrophysiological data. Thus, gross cerebellar damage was moderate, as noted by other authors. According to Okinaka et al7 the extensive cortical damage might also contribute to producing ataxia. The gross phonation and deglutition in case 1 might have been the result not only of the bilateral pyramidal lesions but also of demyelination in the ninth and tenth cranial nerves. We have been unable to find previous reports of the latter. It is only experimentally in animals with induced poisoning by methyl mercury compounds that lesions could be produced in the dorsal roots of the spinal nerves.2 8 9

**Table 3** Nerve conduction velocity

<table>
<thead>
<tr>
<th>Nerve conduction velocity</th>
<th>Immediately after admission</th>
<th>Six months later</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motor conduction velocity</td>
<td>mis</td>
<td>mis</td>
</tr>
<tr>
<td>Median nerve (elbow to wrist)</td>
<td>60</td>
<td>62</td>
</tr>
<tr>
<td>Ulnar nerve (elbow to wrist)</td>
<td>55</td>
<td>60</td>
</tr>
<tr>
<td>Peroneal nerve (knee to ankle)</td>
<td>47</td>
<td>55</td>
</tr>
<tr>
<td>Sensory conduction velocity</td>
<td>mis</td>
<td>mis</td>
</tr>
<tr>
<td>Median nerve (elbow to wrist)</td>
<td>38</td>
<td>59</td>
</tr>
<tr>
<td>Ulnar nerve (elbow to wrist)</td>
<td>26</td>
<td>56</td>
</tr>
</tbody>
</table>

**Table 4** Mercury in nervous system (wet tissue; μg/g; maximum permitted 0.6 μg/g)

<table>
<thead>
<tr>
<th>Occipital cortex</th>
<th>Cerebellar cortex</th>
<th>Frontal cortex</th>
<th>Lenticular nucleus</th>
<th>Thalamus</th>
<th>Optic nerve</th>
<th>Centrum ovale</th>
<th>Choroid plexuses</th>
<th>Pons</th>
<th>Spinal cord</th>
<th>Callosum</th>
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<tbody>
<tr>
<td>Case 1</td>
<td>-</td>
<td>22</td>
<td>10.6</td>
<td>10.6</td>
<td>7.2</td>
<td>7.0</td>
<td>6.6</td>
<td>6.3</td>
<td>4.3</td>
<td>3.5</td>
</tr>
<tr>
<td>Case 2</td>
<td>21.3</td>
<td>13.9</td>
<td>10.6</td>
<td>10.6</td>
<td>7.2</td>
<td>7.0</td>
<td>6.6</td>
<td>6.3</td>
<td>4.3</td>
<td>3.5</td>
</tr>
</tbody>
</table>

**Table 5** Mercury in viscera (wet tissue; μg/g)

<table>
<thead>
<tr>
<th>Kidney</th>
<th>Adrenal</th>
<th>Liver</th>
<th>Stomach</th>
<th>Lung</th>
<th>Spleen</th>
<th>Myocardium</th>
<th>Skeletal muscle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>28.6</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Case 2</td>
<td>84.0</td>
<td>28.3</td>
<td>28.2</td>
<td>23.3</td>
<td>18.2</td>
<td>10.0</td>
<td>6.7</td>
</tr>
</tbody>
</table>
Sensory disturbances have also been disclosed by the slowing down of sensory nerve conduction in cases 3 and 4 found on admission which became normal six months later on clinical recovery (table 3). Electrophysiological data have hitherto invalidated the hypothesis of peripheral neuropathy produced by methyl mercury poisoning.\textsuperscript{10}

Wasting of muscles and twitching in cases 1 and 2 were suggestive of spinal cord anterior horn lesions, which were confirmed by the anatomical evidence. Poisoning by methyl mercury compounds has only very infrequently been noted to produce spinal lesions and these usually spare the ventral horn neurones.\textsuperscript{11} Yet in cases of poisoning with other organic mercury compounds, appearances like amyotrophic lateral sclerosis have been noted, as for example in ethyl phenyl mercury poisoning\textsuperscript{12} and in poisoning with ethyl mercury compounds.\textsuperscript{13} Okinaka et al\textsuperscript{17} consider that organic mercury compounds, according to the chemical nature of the substance, produce two forms of poisoning, a cerebro-cerebellar and a spinal form.

Apart from the lesions located in the cerebral cortex, striate nuclei, cerebellum, and mesencephalon reported by numerous authors\textsuperscript{4,7,14-18}, we noticed some as yet unreported lesions. These were the neuronal lesions of the bulbar reticular formation found in case 1 which we have not found reported before, though certain modifications in the lower brainstem were suspected to occur in organic mercury poisoning by von Burg and Rustam\textsuperscript{19} in their electrophysiological studies. We also failed to find any report of anatomical damage to skeletal muscle or myocardium. It should be noted that, as we noted in our cases, electrocardiographic alterations are associated with ethyl mercury\textsuperscript{14,10} much more frequently than with methyl mercury poisoning.

Toxicological determinations on our cases showed greatly raised levels of mercury in the blood, hair, and urine (tables 1 and 2). The urinary excretion appears to be a less reliable index of the severity of the poisoning, since the highest excretion of mercury was in case 3 where the course was favourable, but urinary excretion of mercury is known to be extremely variable.\textsuperscript{20-22} The levels of mercury in the blood and hair are the most reliable indices of the degree of poisoning, and they were very much increased in case 2 which had a fatal course.

The brain, which is the crucial organ in intoxication by organic mercury compounds, also showed greatly increased levels of mercury in our two necropsy cases. In case 2, when samples were taken from various regions of the central nervous system, the highest levels of mercury were found in the occipital cortex, the cerebellar cortex, the lenticular nucleus, and the frontal cortex (table 4), which were also the regions with evident anatomical lesions.

In case 4 where treatment with penicillamine was begun, urinary excretion of mercury increased and then slightly decreased, but no conclusion can be drawn concerning the effectiveness of such therapy, as case 3 also improved without it.

The published data on treatment with chelating substances in organic mercury poisoning are not conclusive either, as certain authors have failed to obtain positive effects\textsuperscript{13,23} while others reported having obtained improvements.\textsuperscript{5,22} According to Berlin and Ulberg\textsuperscript{24} accumulation of mercury in the brain is accelerated by experimental administration of dimercaprol (BAL).

Indirect poisoning after eating the meat of hogs fed by mistake with seed treated with methyl mercury compounds has also been reported by some American authors.\textsuperscript{25-27}

From study of our cases, it appears that ethyl mercury compounds display a very high toxicity not only for the brain, but also for the spinal motoneurones, peripheral nerves, skeletal muscles, and myocardium.

References

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