Pathology of deaths associated with “ecstasy” and “eve” misuse

C M Milroy, J C Clark, A R W Forrest

Abstract

Aims—To study the postmortem pathology associated with ring substituted amphetamine (amphetamine derivatives) misuse.

Methods—The postmortem findings in deaths associated with the ring substituted amphetamines 3,4-methylenedioxymethylamphetamine (MDMA, ecstasy) and 3,4-methylenedioxymethylamphetamine (MDA, eve) were studied in seven young white men aged between 20 and 25 years.

Results—Striking changes were identified in the liver, which varied from foci of individual cell necrosis to centrilobular necrosis. In one case there was massive hepatic necrosis. Changes consistent with catecholamine induced myocardial damage were seen in five cases. In the brain perivascular haemorrhagic and hypoxic changes were identified in four cases. Overall, the changes in four cases were the same as those reported in heat stroke, although only two cases had a documented history of hyperthermia. Of these four cases, all had changes in their liver, three had changes in their brains, and three in their heart. Of the other three cases, one man died of fulminant liver failure, one of water intoxication and one probably from a cardiac arrhythmia associated with myocardial fibrosis.

Conclusions—These data suggest that there is more than one mechanism of damage in ring substituted amphetamine misuse, injury being caused by hyperthermia in some cases, but with ring substituted amphetamines also possibly having a toxic effect on the liver and other organs in the absence of hyperthermia.

Keywords: eve, ecstasy, postmortem pathology

Recreational use of 3,4-methylenedioxymethylamphetamine (MDMA), more commonly known as “ecstasy” (and a variety of other names including “XTC”, “Adam” or “E”), is now well established. In Britain upwards of 500,000 people are said to use the drug each week (Harris Poll (1992) for “Reportage”, BBC2, 22 Jan 1993).

MDMA is a ring substituted amphetamine with psychoactive properties. First synthesised in 1914 from methyleneoxyamphetamine (MDA), itself a drug of misuse (known as the “love drug”), it has been used in psychotherapy and was originally used as an appetite suppressant. The drug has ceased to be used medicinally and is now an established part of the illegal drug scene. It is banned in most countries. In the UK it is a class A drug as defined in Schedule 2 of the Misuse of Drugs Act 1971. It has no medicinal use in the UK and cannot be prescribed. As well as MDA and MDMA, another variant, methylene-dioxymethylamphetamine (MDA, known as “eve”), which is similarly proscribed, is commonly encountered. All have similar pharmacological effects.

In the UK, MDMA is often taken by young people at discos and rave parties. Both involve dancing, but especially at the latter there is vigorous repetitive dancing in crowded rooms with a hot and humid atmosphere. The dangers of this activity are recognised to a certain extent as rooms to “chill out” are often available for people to rest in after periods of exertion. Toxic effects and the occasional death following ring substituted amphetamine misuse have been reported but postmortem data are lacking.1-20 In this paper we report on deaths associated with ring substituted amphetamine misuse and detail the postmortem findings.

Methods

Seven deaths have been investigated by the University of Sheffield Department of Forensic Pathology in the past three years, which were associated with ring substituted amphetamine misuse. One case has been reported previously.26 Case details are presented in table 1. All of the subjects were white men, between 20 and 25 years of age. Three of the victims collapsed at a rave or disco, two were found in bed, one in a collapsed state and one dead, one collapsed in the street, and one was admitted to hospital with progressive jaundice.

Only two of the men had documented hyperthermia, with temperatures of 44°C and 39.5°C. One of the deaths was thought to have been caused by water intoxication, the victim having drunk an estimated 14 litres of water in trying to quench the thirst that is frequently associated with ingestion of ring substituted amphetamines. None of the men were known intravenous drug abusers.

Full drug screening was performed on body fluids, including a screen for drugs of abuse. Screening by gas chromatography/mass spectrometry was also performed.

Results

Pathology

The pathological and toxicological data are summarised in table 1.

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References


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Table 1  Clinical features, toxicology and postmortem findings

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Age (years)</th>
<th>Clinical data</th>
<th>Toxicology (blood concentrations)</th>
<th>Pathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>21</td>
<td>Collapsed at rave. Agitation, unconscious, 44°C, cardiac arrest</td>
<td>MDMA (4-2 mg/l) Amphetamine (1-4 mg/l)</td>
<td>Heart—contraction band necrosis Liver—focal necrosis Brain—focal haemorrhage, neuronal degeneration</td>
</tr>
<tr>
<td>2</td>
<td>20</td>
<td>Collapsed at disco. Thirst, convulsions, 36°C, TPR low Na+, unconscious. Water intoxication</td>
<td>MDMA (0-04 mg/l)</td>
<td>Heart— widespread foci of necrosis Liver—foci of necrosis Brain—gross oedema, focal haemorrhage.</td>
</tr>
<tr>
<td>3</td>
<td>24</td>
<td>Collapsed and died at disco. ? Temperature Found dead in bed after party. No symptoms</td>
<td>MDEA (0-187 mg/l) Amphetamine (0-453 mg/l)</td>
<td>Heart—focal necrosis Liver—focal necrosis Brain—normal</td>
</tr>
<tr>
<td>4</td>
<td>21</td>
<td>Temperature Found dead in bed after party. No symptoms</td>
<td>MDMA (2.1 mg/l) MDEA (3-5 mg/l)</td>
<td>Heart—focal necrosis Liver—focal necrosis Brain—focal haemorrhage Inhalation of vomit</td>
</tr>
<tr>
<td>5</td>
<td>20</td>
<td>Found unconscious in bed. Rigidity, 34-5°C, TPR, low Na+, cerebral hypoxia. Survived four days</td>
<td>MDMA (0-09 mg/l) MDA (0-13 mg/l) Amphetamine (0-256 mg/l)</td>
<td>Heart—widespread foci of necrosis Liver—extensive necrosis Brain—hypoxic changes, DIC Lung—pulmonary infarcts Heart—widespread foci of necrosis Liver—focal necrosis</td>
</tr>
<tr>
<td>6</td>
<td>25</td>
<td>Sudden collapse in street. Previously well, witnessed collapse. Brought in dead</td>
<td>MDMA (trace in urine) MDA (trace in urine)</td>
<td>Heart—normal Liver—massive hepatic necrosis Brain—normal</td>
</tr>
<tr>
<td>7</td>
<td>23</td>
<td>Progressive jaundice and liver failure. All investigations negative. Admitted heavy ecstasy use, even after onset of jaundice</td>
<td>Admitted regular ecstasy use</td>
<td></td>
</tr>
</tbody>
</table>

LFTs = liver function tests.

Liver
The liver showed dramatic changes. Necrosis was seen in all cases. In one case there was centrilobular and midzonal necrosis (fig 1). This patient survived for four days. In the other cases focal necrosis in zone 3 was present, with an acute inflammatory response surrounding necrotic hepatocytes (fig 2). Fatty change, sinusoidal dilatation and inflammation were also identified. No other hepatic pathology was identified in these cases. Postmortem virology was not done in these cases. In case 7 there was massive hepatic necrosis, with the liver only weighing 795 g at necropsy. Full viral screening was negative in this case. The livers were of normal size in the other cases.

Heart
Changes were seen in five of the seven hearts examined. Histologically, the changes ranged from contraction band necrosis to individual myocyte necrosis with a surrounding neutrophil and macrophage inflammatory response; the same changes as have been described in catecholamine induced myocardial injury (fig 3). In one case foci of fibrosis were identified in the heart of a man in whom traces of MDMA and MDA were found in the urine.

Brain
The brain of one of the cases of rapid death showed disseminated intravascular coagulation (DIC), oedema and degeneration of neurones, particularly apparent in the locus ceruleus. Two cases showed foci of haemorrhage (fig 4). One case showed severe cerebral oedema consistent with water intoxication and had additional occasional perivascular haemorrhages.

Other organs
Pulmonary infarction was seen in case 5. In two of the other rapid deaths intra-alveolar haemorrhage was present. In one case there was inhalation of gastric contents. In the patient who died of water intoxication there was complete necrosis of the pituitary gland. Severe cerebral oedema was present and the pituitary necrosis was probably a direct consequence of this as the blood supply was compromised. In two cases the kidneys were examined for myoglobin, which was not detected. No skeletal muscle damage was identified.

TOXICOLOGY
The toxicological data are shown in table 1. MDMA was found in five cases, varying from trace concentrations to 4-2 mg/l. MDEA was found in two cases, at concentrations of 3-5 and 0-187 mg/l. Amphetamines were detected in three cases at concentrations of 0-256, 0-453

![Figure 1  Section of liver showing centrilobular and midzonal necrosis following ingestion of "ecstasy".](attachment:image)
sensations at raves and discos. However, when 29 volunteers took MDMA, as well as pleasant
symptoms, all complained of undesirable
effects which included loss of appetite, trismus
and bruxism, nausea, muscle aches, stiffness,
and ataxia.\textsuperscript{27} Sweating, tachycardia and hy-
pertension, insomnia, and fatigue were also
reported. More serious complications reported
include hyperthermia, convulsions, other
cardiac arrhythmias, rhabdomyolysis, dis-
seminated intravascular coagulation, renal fail-
ure, hyponatraemia, hepatotoxicity, pneumo-
mediastinum, aplastic anaemia, cerebral
infarction, cerebral haemorrhage, and cerebral
venous sinus thrombosis.\textsuperscript{1-26}

In five of the cases reported here com-
lications followed shortly after ingestion. Four
of these deaths seem to be linked directly to
the toxic effects of MDMA and MDEA. In the
fifth case there was evidence of water in-
toxicaiton, which followed the taking of “ec-
stasy”. In this case, however, myocardial and
liver pathology was identified which was similar
to the other cases. The sixth case collapsed in
the street. Myocardial fibrosis and foci of liver
necrosis were identified and toxidology revealed
traces of MDMA.

MDMA is believed to act on at least three
neurotransmitter pathways, as does cocaine,
but with MDMA the serotonergic (5-hydroxy-
tryptamine) pathway is principally affected,
which would account for the more pronounced
effect on mood. Cocaine acts chiefly on the
dopaminergic system, which accounts for its
greater addictive properties. MDMA also acts
on the noradrenergic system. Serotonin plays a
major role in thermoregulation and interference
with this mechanism is believed to be the cause
of the hyperthermia which arises as a com-
plication of ring substituted amphetamine mis-
use. Stimulation of the noradrenergic system
also probably contributes to hyperthermia.

Hyperthermia may account for many of the
changes seen in deaths from ring substituted
amphetamine misuse, although it is interesting
to note that raised temperatures were only
documented in two of our cases.

The pathology of heatstroke has been re-
ported in a number of papers.\textsuperscript{28-34} In the liver
the most striking change is centrilobular
necrosis. Sinusoidal congestion and dilatation, and
portal and sinusoidal inflammation may be
present. Fatty change has been reported oc-
casionally. Cholestasis may be present, espe-
cially in fatal cases. Rubel and Ishak\textsuperscript{34} did
not find liver necrosis as frequently as other
authors. They examined the liver in 50 military
recruits who had died of heatstroke. These men
were predominantly white. Kew \textit{et al}\textsuperscript{41} had
found liver necrosis a common finding in black
South African gold miners. The difference in
frequency may be related to the fact that the
gold miners work in very high environmental
temperatures with over 90\% humidity, con-
ditions not dissimilar to some raves and discos.

In the myocardium contraction band nec-
rosis and small foci of necrosis with a mixed
inflammatory infiltrate are seen. These features
are also evident in catecholamine induced in-
jury. In the brain ring haemorrhages and hyp-
oxic changes have been described. The kidneys may show acute tubular necrosis and pulmonary haemorrhage is common.

The mechanism of damage in heatstroke is postulated to be caused by circulatory collapse and hypoxic damage, possibly combined with disseminated intravascular coagulopathy, which has been recorded in heatstroke,\textsuperscript{36,37} and as a complication of MDMA and amphetamine ingestion.\textsuperscript{37} In the myocardium the damage may be related to catecholamine induced injury. These myocardial changes are also seen secondary to brain pathology, and contraction band necrosis may be seen in resuscitation, especially when catecholamines are used. Whilst these are alternative explanations for the changes seen, not everyone was resuscitated, and the myocardial changes are frequently seen, even in early deaths.

The pathological changes present in these deaths are the same as those seen in deaths from heatstroke. These changes provide further evidence that hyperthermia can cause death following misuse of ring substituted amphetamines. Evidence of disseminated intravascular coagulopathy was also present in the brain. These deaths may therefore be a complication of hyperthermia, DIC and shock. Some caution must be exercised, however, in ascribing a death from ring substituted amphetamine misuse to these mechanisms as a high temperature is not recorded in all cases, although hyperpyrexia may have been present at some time.

In the seventh case death was caused by fulminant liver failure and postmortem examination revealed massive hepatic necrosis. All investigations for the cause of liver failure were negative. The young man admitted to regular and heavy ecstasy use, and this case is similar to cases described by Henry et al.\textsuperscript{7} There is increasing evidence that “ecstasy” is hepatotoxic, and liver changes have been reported following biopsy.\textsuperscript{10,11} These changes, however, differed from those found in the acute deaths described above. Whether the damage is caused by an idiosyncratic reaction to MDMA, or a contaminant of the drug is unclear. Therefore, there seems to be a second mechanism for liver injury from ring substituted amphetamines unrelated to hyperthermia. Furthermore, the possibility that the myocardium may also be damaged by these drugs without documented hyperthermia is suggested by case 6 and other reports, where myocardial fibrosis has been reported following misuse of MDMA.\textsuperscript{3}

Individual susceptibility to ring substituted amphetamines may be related to its metabolism

<table>
<thead>
<tr>
<th>Street name</th>
<th>Appearance</th>
<th>Size (mm)</th>
<th>Weight (mg)</th>
<th>Active drug (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adam &amp; Eve</td>
<td>Round white tablet, E/A on one surface</td>
<td>12 x 2</td>
<td>380</td>
<td>MDEA (56)</td>
</tr>
<tr>
<td>California Sunrise</td>
<td>Round, scored off white tablet</td>
<td>11 x 4</td>
<td>410</td>
<td>Amphetamine (22)</td>
</tr>
<tr>
<td>Green Burger</td>
<td>Round grey green speckled tablet</td>
<td>10 x 2</td>
<td>300</td>
<td>Amphetamine (6-2)</td>
</tr>
<tr>
<td>Love Heart</td>
<td>Round white tablet, heart on back</td>
<td>8 x 3</td>
<td>120</td>
<td>Amphetamine (6-2)</td>
</tr>
<tr>
<td>MDMA Capsule</td>
<td>White capsule</td>
<td>5 x 17</td>
<td>190</td>
<td>Pseudoephedrine</td>
</tr>
<tr>
<td>Power Pack</td>
<td>Biconvex, scored white tablet</td>
<td>8 x 5</td>
<td>300</td>
<td>MDEA (32)</td>
</tr>
<tr>
<td>Red Devil</td>
<td>Round tablet, pink and white with grey speckles</td>
<td>10 x 2</td>
<td>340</td>
<td>MDEA (0-16)</td>
</tr>
<tr>
<td>Snowball Split</td>
<td>Biconvex white tablet</td>
<td>8 x 4</td>
<td>390</td>
<td>Pseudoephedrine</td>
</tr>
<tr>
<td></td>
<td>Round white scored tablet</td>
<td>8 x 2</td>
<td>200</td>
<td>MDEA (0-16)</td>
</tr>
<tr>
<td>Triple X</td>
<td>Round scored tablet, white with black speckles</td>
<td>8 x 4</td>
<td>260</td>
<td>Pseudoephedrine</td>
</tr>
<tr>
<td>White Burger Saucer</td>
<td>Round grey &amp; white speckled tablet</td>
<td>11 x 2</td>
<td>390</td>
<td>MDEA (57-2)</td>
</tr>
<tr>
<td>White Cally</td>
<td>Round white tablet</td>
<td>8 x 2</td>
<td>190</td>
<td>MDEA (59-21)</td>
</tr>
<tr>
<td>White Cap</td>
<td>Dirty white capsule</td>
<td>15 x 5</td>
<td>280</td>
<td>Pseudoephedrine</td>
</tr>
</tbody>
</table>

Figure 4 Perivascular haemorrhage in the cerebral cortex following ingestion of “ecstasy”.
in the liver. Abnormalities have been reported in demethylation in susceptible individuals, related to human debrisoquine hydroxylase (CYP2D6). Another problem for the recreational drug taker is the quality of the tablets they take. Examination of "ecstasy" tablets by one of the authors (ARWF) has shown that the contents of the tablets vary greatly, and may be a mixture of MDA, MDEA, amphetamine, or no pharmacologically active substance; compounds such as potassium chloride may also be found (table 2). With such material the possibility of toxic contaminants being present is evident.

The short term risks of "ecstasy" use are becoming increasingly more apparent and questions must be asked about the long term effects on the brain, liver and heart considering the pathology found during autopsies in those who die.

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