Myocardial infarction: a complication of amitriptyline overdose

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Summary: A 22 year old woman was admitted with amitriptyline overdose. Twenty six hours later she developed acute myocardial infarction. Cardiotoxic effects of tricyclic antidepressants are discussed and the importance of considering myocardial infarction as a complication of tricyclic antidepressant overdose is emphasized.

Introduction

Tricyclic antidepressants (TCA) overdose represents 25% of serious drug self-poisoning, with an overall mortality rate of up to 10%. Evidence of cardiovascular toxicity is present in the majority of TCA overdoses. Although myocardial infarction has been reported in patients receiving therapeutic doses of TCA, it has not, to our knowledge, been reported following TCA overdose. We describe such a case in a young patient.

Case Report

A 22 year old woman was admitted 24 hours after taking 300 mg of amitriptyline and 80 mg of diazepam. Apart from a peptic ulcer she had no history of diabetes or any other illnesses, and was not on any medication. She had never smoked and there was no family history of ischaemic heart disease.

On admission she was unconscious and restless, the pulse was 130 beats per minute and her blood pressure was 110/70 mm Hg. The heart sounds were normal and she had a soft systolic murmur at the pulmonary area. The pupils were dilated but reactive to light. Noneurological localizing signs were noted. The electrocardiogram (ECG) revealed sinus tachycardia and the QRS duration was 84 ms. Two hours later her blood pressure fell to 90/60 mm Hg and ST segment elevation was noted. A 12 lead electrocardiogram showed changes of an acute anteroseptal myocardial infarction (Figure 1). The PR, QRS and QTc duration remained normal. Serial measurements of creatine kinase (CK), creatine kinase M.B., aspartate aminotransferase (AST) and lactic dehydrogenase (LDH) revealed a classical pattern consistent with acute myocardial infarction (Table I). Two dimensional echocardiography was performed and revealed hypokinesia of the septum. The valves appeared normal and the heart chambers were not dilated.

She regained consciousness 12 hours later but she did not recall having chest pain. She did not have fever or a pericardial rub or gallop rhythm at any stage during admission. Her platelets were normal. The blood sugar was 4.5 mmol/l and total cholesterol was 4.7 mmol/l. The anti-phospholipid antibody test was not performed. She made an uneventful recovery and was discharged home well. She failed to attend for follow-up.

Discussion

Cardiovascular effects of TCA overdose are common. The most common abnormalities are sinus tachycardia, prolongation of the PR, QRS and QTc intervals, bundle branch block, arrhythmias, hypotension and cardiac arrest. These effects are mediated by the anticholinergic properties of the TCA, their ability to potentiate the action of the noradrenaline (which could elicit arrhythmias), their blocking effects of peripheral alpha adrenergic receptors and by their quinidine-like action. The latter two mechanisms can contribute to recumbent hypotension sometimes seen in TCA overdose. Similar to quinidine TCA can cause impaired intracellular conduction and depression of myocardial

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Table 1 Cardiac enzymes changes consistent with acute myocardial infarction.

<table>
<thead>
<tr>
<th>Day</th>
<th>CK (normal 24-170 IU/l)</th>
<th>CKMB (normal &lt;8 IU/l)</th>
<th>AST (normal 5-45 IU/l)</th>
<th>LDH (normal 100-225 IU/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>490</td>
<td>38</td>
<td>258</td>
<td>351</td>
</tr>
<tr>
<td>2</td>
<td>630</td>
<td>54</td>
<td>325</td>
<td>394</td>
</tr>
<tr>
<td>3</td>
<td>110</td>
<td>2</td>
<td>267</td>
<td>333</td>
</tr>
</tbody>
</table>

Day 1 sample was taken 5 hours after admission. Day 2 and 3 samples were performed at 24 and 48 hours later respectively.

contractility. ST segment elevation resembling that of myocardial infarction has been described following TCA overdose. Marshall et al. reviewed four studies including 289 patients who had TCA overdose and no cases of myocardial infarction were found. Cardiac arrest has been reported in up to 12% of patients in one of these studies and no post-mortem findings were given. TCA poisoning outcome is unpredictable since severe cardiac complications including cardiac arrest have been reported in patients who ingest doses as low as 200 mg of imipramine.

Our patient took only 300 mg of amitriptyline and yet developed acute changes of myocardial infarction 26 hours later. She was unconscious at the time she developed the changes but serial electrocardiograms and cardiac enzymes confirmed the diagnosis. The exact pathogenic mechanism by
which myocardial infarction occurred in our patient remains uncertain. However, several hypotheses could be postulated.

Coronary vasospasm can cause myocardial infarction in some patients with normal coronary arteries. However, TCA are not known to produce any vasospastic changes in coronary arteries. Animal studies have shown that small doses of TCA given intravenously tend to increase blood pressure, cardiac contractility and coronary artery blood flow, whilst large doses causes a decrease in these parameters. When this animal data is extrapolated to humans, the dose at which the cardiovascular depressant effects begin to predominate would be about equivalent to a high daily therapeutic dose.

Hypotension occurs in 26–65% of patients with TCA overdose, and is usually secondary to myocardial depression and/or peripheral alpha adrenergic blockade. Significant hypotension (systolic blood pressure under 100 mm Hg) was observed at some point during the first 24 hours in 53% of patients with amitriptyline overdose. Hypotension which can precipitate a myocardial infarction seems to be the likely explanation in our patient.

Surveillance of patients taking TCA overdose with serial cardiac enzymes and electrocardiograms may reveal more cases of myocardial infarction complicating this type of overdose.

Acknowledgements

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References

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