A Case of Severe Amitriptyline Intoxication: Electrocardiography Signs Suggesting Ventricular Tachycardia
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ABSTRACT

Amitriptyline poisoning may result in severe neurological and cardiac problems. We report a case of (13-month girl) severe amitriptyline intoxication with neurological, respiratory and electrocardiographic findings. With respiratory support, seizure control, NaHCO₃ infusion, intravenous lipid infusion, selective plasma exchange and arrhythmia control the patient recovered. We stressed that amitriptyline intoxication may lead to severe problems. Electrocardiographic findings may not be differentiated from ventricular tachycardia. In these patients, survey may be well with aggressive treatment modalities.

Keywords: Aggressive treatment, amitriptyline, intoxication, ventricular tachycardia

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INTRODUCTION

Amitriptyline is one of the most commonly used tricyclic antidepressants. Clinical results of amitriptyline poisoning are well defined in adults and in children (1–4). Amitriptilin poisoning may result in severe neurological and cardiac problems. Treatment may be challenging and even death may occur (5). In this paper we aimed to report a child with severe amitriptyline poisoning and discuss the urgent treatment options.

CASE REPORT

A 13-month old girl was admitted to emergency department of the local state hospital with the suspicion of drug intake (Laroxyl® 10 mg drage, containing amitriptyline) 2 hours previous to the admission. The amount of the possibly ingested drug was not known. The patient had been admitted after recognition of the newly developed somnolance. Physical findings had revealed unconsciousness, flexor response to painfull stimuli and bilateral negative pupillary light reflex. After gastric lavage, active choarchal had been administered. Complete blood count, liver and renal functions and serum electrolytes had found to be normal. Blood gase analysis had found as following; pH=7.24, pCO₂=54mmHg, HCO₃=19.4mmol/L, Lac=2.7mmol/L. During monitorization ventricular tachycardia (VT) (Figure- 1 and Figure-2) had been detected and lidocain infusion had been started. Tonic clonic convolution had been controlled with intravenous diazepam infusion. NaHCO₃ infusion (2 mEq/kg dose in 20 minutes) had been started and the patient transported to our pediatric emergency department.
Fig. 1: The first electrocardiography that was evaluated as ventricular tachycardia (2nd hour of the event).

Fig. 2: Frequent ventricular premature contractions with sinus beats. Configuration of the premature contractions are not identical with the QRS morphology of the first electrocardiography.

The patient admitted to our emergency department at the 8th hour of the event.

The patient was unconscious, the Glasgow Coma Score (GCS) was 4 point, pupils were fixed dilated and pupillary light reflex was negative bilaterally. The patient had shallow breathing. Rhythm was irregular (Figure-3), heart rate was 140/bpm and blood pressure was 75/41 mmHg. During the first examination generalized tonic clonic convulsion developed and was controlled by intravenous diazepam injection. She was intubated. Midazolam infusion was started. Intravenous fluid administration, containing NaHCO₃ was started with a rate of 2000 ml/m². Dopamin infusion was started because of low blood pressure.
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The first electrocardiography (ECG) recordings (from the first admission center) revealed a wide QRS tachycardia (Figure- 1) with a right superior axis deviation (QRS duration was 0.15 s, QTc was measured as 0.47-0.60 s). “P” waves could not be identified. In another ECG (Figure- 2) frequent ventricular premature contractions with sinus beats were present. Configuration of the premature contractions was not identical with the QRS morphology of the first ECG. The first ECG recorded in our emergency room showed a wide QRS tachycardia (140 bpm) with a right bundle branch block and right inferior axis deviation and intermittent sinus beats after a short pause (Figure- 3). So, a definite differentiation between VT and a wide QRS sinus tachycardia as result of amitriptyline overdose (4, 6, 7) could not be done. Lidocain infusion was continued.

![ECG recordings](image)

**Fig. 3:** The first electrocardiography recorded in our emergency room showed a wide QRS tachycardia (140 bpm) with a right bundle branch block and right inferior axis deviation with intermittent sinus beats (arrow) after a short pause (8th hour of the event).

Two additional active charcoal was given with a 4-hour interval. Intravenous 20% lipid solution was administered (1.5 ml/kg bolus and 0.25 ml/kg in 30 minutes). Echocardiographic evaluation revealed a normal heart with normal left ventricular functions. Complete blood count, liver function tests, renal functions and serum electrolyte levels (Na+=138 mmol/L, Ca=8.8 mg/dL, K=3.5 mmol/L) were normal. Blood gase analysis showed respiratory acidosis (pH=7.26 pCO2=57.4 mmHg, HCO3=25.1 mmol/L, Lac=1.6 mmol/L,
BE=-1) but returned to normal at the first hour of mechanical ventilation (pH=7.35, pCO2=44.2 mmHg, HCO3=24.1 mmol/L, Lac=2.9 mmol/L, BE=-0.7).

Despite the continuing NaHCO3 infusion, pH value was still not upper than 7.35 and the ECG findings persisted. So, selective plasma exchange was started at the 15th hour of drug intake. During the procedure dobutamine was added to the treatment because of persisting low blood pressure. Selective plasma exchange was stopped at the 7th hour of the procedure. Hemoglobin level decreased to 7.3 g/dl, so erythrocyte suspension was administered.

During this course, QRS morphology remained stable but the QRS duration become shorter and shorter (Figure- 4). In the ECG obtained at the 10th hour of event “p” waves begin to appear, so lidocaine was ceased. At the 20th hour of event the sinus rhythm with a right axis deviation settled. The PR interval was 0.14 s and the QRS duration was 0.08 s (Figure- 4). At the 27th hour of event, right axis deviation was also corrected (Figure- 5). All these findings suggested that the ECG findings were the result of amitriptyline overdose, not VT.
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Fig. 4: The electrocardiography at the 9th (A), 10th (B) and 20th (C) hours of event. QRS morphology remained stable but the QRS duration become shorter and shorter. In the electrocardiography obtained at the 10th hour (B), “p” waves begin to appear, and it was apparent in the 20th hour electrocardiography (C).

Fig. 5: At the 27th hour of event, right axis deviation was also corrected.

After completion of the plasma exchange, sedation was stopped and the patient was extubated successfully at approximately 32th hour of event (GCS=14 point). Dopamin, dobutamine and NaHCO3 infusions were stopped. The patient went to normal room at the 34th hour of event.
DISCUSSION

Amitriptylin is one of the most common used antidepressants. Also the tricyclic antidepressants, especially the amitriptyline, are the most common used drugs in drug intoxication worldwide (8, 9). Acute results of amitriptyline intoxication had been well defined both in children and adults (1, 3, 4, 10).

Amitriptyline intoxication yields a variety of clinical signs and symptoms such as tachycardia, hypotension, anticholinergic findings such as dilated pupils, tachycardia, urinary retention, impaired consciousness ranging from lethargy to a comatose state, atrioventricular block, prolonged QTc interval or QRS duration, convulsions and dyspne due to respiratory depression, pulmonary insufficiency and occasionally adult respiratory distress syndrome (6, 7, 11, 12). Life threatening complications like convulsions, arrhythmia and respiratory depression are relatively more frequent among children than in adults (10).

Previous studies have shown that severe neurological and cardiovascular side effects arise from the blockage of voltage-dependent sodium channels (10). Although some studies suggest a relation between the serum drug level and the complications (10), some others indicate that severe toxic effects may occur with lower doses (2).

First-line treatment focuses on general measures as well as correction of electrolyte disorders and arrhythmia (6, 9). If the patient admits during the first 12 hours, gastric lavage should be performed and active charcoal should be administered. Respiration should be supported and hydration should be achieved. If conduction disturbance of arrhythmia is observed NaHCO₃ should be given (5). For this purpose administration of hypertonic saline had also been proposed (13).

In case of amitriptyline intoxication, phenytoin is no longer recommended for control of convulsions because of its limited efficacy and possible prodysrhythmic effects. Benzodiazepines are the most commonly preferred anti-convulsant drugs (14). As it is a lipid
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soluble drug intravenous lipid treatment has been applied in patients with amitriptyline intoxication (15, 16).

There is no known antidote against any of the tricyclic antidepressants. The American Society for Apheresis (ASFA) recommends plasmapheresis to support primary treatment in this type of drug poisoning, which does not respond to certain and traditional treatments (17). In patients with life threatening amitriptyline intoxication successful results had been reported with plasma exchange had been (7, 18-21). Recently one study reported successful result with extended high cut-off dialysis (9). In our case, we performed selective plasma exchange due to the persisting life threatening clinical condition despite all other treatment options.

Paksu et al (10) reported a relation between the serum sodium level and ECG findings and complications. However, in our patient despite the severe ECG findings neurological signs and depressed respiration the serum sodium level was normal.

The cardiac effects of tricyclic antidepressant poisoning are caused by a blockage of cardiac sodium channels in the His- Purkinje system and ventricular muscle. The ECG manifestations of slowed cardiac depolarization are prolongation of the QRS complex and an R-wave amplitude of 3 mm or higher in lead aVR (22). Some severe arrhythmias may also occur in these patients. Streanga et al (12) reported occurrence of ventricular premature beats, isolated, couplets and triplets, VT and torsade de pointes, and severe ventricular repolarization disturbances with diffuse subendocardial ischemia in 8 children with signs of amitriptyline intoxication.

Anticholinergic effects of the drug results in tachycardia (5). Slowing of intraventricular conduction, manifested by prolonged PR, QRS and QT intervals on the standard ECG are also well defined surface ECG findings of toxicity of tricyclic antidepressants (23). Wide QRS with prolonged PR and QT intervals results in disappearance “p” waves in the previous T waves. Because of this finding, the rhythm may be misdiagnosed as VT and cardioversion may be applied, especially in patients in whom rhythm is
monitorised from one lead. So, in these patients 12-lead surface ECG should be evaluated and multiple leads should be monitorised for a long period carefully.

In our patient, the rhythm had been thought as VT (Figure-1) in the first admitted center and lidocaine infusion had been started. In the second ECG (Figure-2) there was frequent ventricular premature beats and the configuration of ventricular premature beats was not identical with the QRS morphology of the first ECG. Although we thought the first ECG as a result of amitryptiline toxicity, differentiation of VT could not be made. In the first ECG obtained in our clinic intermittent sinus beats were seen during a wide QRS tachycardia (Figure-3). During follow-up heart rate gradually decreased, QRS duration shortened but QRS morphology remained similar (Figure-4), and approximately in the ECG obtained at the 10th and 20th hours of event (Figure-4) “p” waves became apparent. With appearance of “p” waves VT was eliminated. These findings suggested us that the previous ECG findings were also result of amitriptyline effect not VT. At discharge, ECG was normal for the age of the patient (Figure-5).

Amitriptyline intoxication may lead to severe neurological, respiratory and cardiac problems. ECG findings may not be differentiated from VT. In these patients survey may be well with aggressive treatment modalities. In addition to standard treatments, plasmapheresis should be kept in mind.
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