Clinical Findings of Emergence After Olanzapine Overdosage: Pediatric Case Report and Review of Literature

Olanzapin Aşırı Alımı Sonrasında Ortaya Çıkan Klinik Bulgular: Çocuk Olgu Sunumu ve Literatürün Gözden Geçirilmesi

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ABSTRACT

Introduction: Olanzapine is an atypical antipsychotic drug of the thienobenzodiazepine group; its clinical use has increased in the last 2 decades. Especially in pediatric cases, the clinical experience in overdose of olanzapine and other atypical antipsychotics is limited. Overdosage of atypical antipsychotic drugs, including olanzapine, predominately causes central nervous system depression and anticholinergic effects.

Case Report: We present the case report of a 3-year-old female child who ingested 20 mg of olanzapine, which resulted in clinical findings, such as mental status changes, peripheral edema and rapid atrial rhythm, temporary prolonged P-R interval, and T-wave inversion.

Conclusion: According to the current literature, olanzapine seems to be a medium-level safe atypical antipsychotic drug in terms of clinical use. Therefore, the pediatric cases should be monitored closely with accidental olanzapine overdosage.

Keywords: Olanzapine, overdosage, pediatric case

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Introduction

Having been marketed for almost 17 years as an atypical antipsychotic drug, olanzapine is a new atypical antipsychotic drug that has become widespread in prescriptions day by day. Olanzapine is used for the treatment of psychoses, schizophrenia, schizoaffective disorders, bipolar disorders, and other conditions with psychotic or delusional components (1). Presumably, as the usage of these medications in adults increases, the frequency of unintentional pediatric ingestions will increase.

Evidence on the efficacy and safety of atypical antipsychotics in children and adolescents with schizophrenia is limited. The use of typical antipsychotics has been limited to patients who are resistant to atypical antipsychotics and intolerant to their
adverse effects or require injections or depot preparations. Besides, further double-blind, placebo-controlled trials and long-term safety assessments are needed before definitive conclusions can be reached about the place of atypical antipsychotics in the therapeutic armamentarium of childhood-onset schizophrenia (2).

Olanzapine affects dopaminergic (D1-4), serotonergic (5HT2, 5HT3, 5HT6), histaminergic (H1), muscarinic (M1-5), and adrenergic (a1) receptors. The metabolism of olanzapine is complex; it is primarily metabolized by the cytochrome P-450 system (CYP-450), mostly by the isoenzyme 1A2 system and 2D6. Olanzapine can also cause an increase of serum transaminase levels during medications (3).

Although structurally and functionally related to clozapine, olanzapine possesses a more favorable side effect profile. Although the usual dose range for olanzapine is 5-15 mg/d, there are no standard reference values with respect to the expected concentrations of olanzapine after therapeutic administration. In clinical studies, steady-state blood (plasma) concentrations of olanzapine are rarely over 150 ng/mL, but the potential for toxicity has been suggested at concentrations as low as 100 ng/mL. Generally, new atypical antipsychotic drugs may have a safer therapeutic, side effect, and overdose profile than first-generation antipsychotic drugs. Olanzapine has slight side effects on the extrapyramidal system (EPS) in low dosage intake (4). Olanzapine overdoses in children are generally associated with more significant adverse effects. With another statement, the symptoms in overdose are generally a reflection of olanzapine’s known pharmacological actions and encompass somnolence, mydriasis, blurred vision, respiratory depression, hypotension, and extrapyramidal and anticholinergic effects.

Children who are potentially ingesting a toxic dose or who are symptomatic should be considered for assessment in a hospital. Children, therefore, require more active intervention than adults. Furthermore, they should be monitored closely when ingestion of olanzapine occurs. Most cases resolve with good supportive care (5).

Case Report
A 3-year-old girl was brought to our emergency department on account of accidental digestion of medicine containing olanzapine (Rexapin®) (20-mg tablets; Abdi Ibrahim Medical, Istanbul, Turkey), which is used by her mother.

In the physical examination arrival, the vital parameters were determined as: blood pressure relatively low (systolic=90 mm Hg and diastolic=50 mm Hg), tachycardic (151 bpm), body temperature was normal (36.5°C), respiratory rate: 22/min, and spO₂: 95%. Weight was 15 kg (percentile of range 50%-75%). She was conscious but had a somnolence status with insufficient cooperation. Glasgow Coma Scale (GKS)=10-12. Other system examinations were normal. On the arrival ECG, rapid atrial rhythm was determined after drug intake at the 90th minute (Figure 1). Laboratory examination findings on arrival were: WBC: 7.9 K/μL, Hgb: 14.7 g/dL, PLT: 360 K/μL; glucose: 126 mg/dL, BUN: 20 mg/dL, creatinine: 0.4 mg/dL, AST: 71 U/L, ALT: 101 U/L, LDH: 227 U/L, CK: 81 U/L, Na: 145 mmol/L, K: 4.7 mmol/L, Cl: 9.8 mmol/L, and Ca: 9.4 mg/dL.

After the first physical examination in the emergency department, gastric lavage, activated charcoal (1 g/kg), intravenous fluids, and oxygen (2-3 L/min.) were rapidly applied, and the child case was monitored in supportive care. In this case, treatment in accordance with the symptoms that might occur was planned. On arrival, for the hypotension and tachycardia, 1500 cc ringer lactate was applied (for the first 24 hours) as intravenous fluid. In the following days, she was given approximately 1000 cc ringer lactate as intravenous fluid every 24 hours. No medication applications were needed. At the 5th hour of observation, sinus rhythm and prolonged P-R interval were determined as 103 bpm after drug intake at the 5th hour (Figure 2), and blood pressure was slightly low (systolic=95 mm Hg and diastolic=55 mm Hg), GCS= 12, and Poisson Severity Score (PSS)=2. However, at the 7th hour of observation, a slight level of sinus tachycardia (117 bpm) and additionally supraventricular premature complexes and T-wave negativities in anterior derivations occurred. Meanwhile, ongoing slightly prolonged P-R interval was determined (Figure 3). After the first night and throughout the entire observation, it was determined that GCS=15. On the 4th day of observation, supraventricular tachycardia (SVT) was monitored, but the slightly prolonged P-R interval recovered itself (Figure 4). On the 10th day, electrocardiogram (ECG) findings were abnormal, except the rapid atrial rhythm recovered (Figure 5). Therefore, on the 14th day, the ECG findings recovered exactly.

The laboratory findings were recorded at the 5th and 8th hours of patient monitoring: AST: 64 U/L and 72 U/L, ALT: 91 U/L and 98 U/L, and LDH: 183 U/L and 192 U/L. CK value progressed at the normal...
limits. Among the electrolytes, the level of sodium decreased down to 135 mmol/L on the first day of observation and 133 mmol/L on the 7th day of observation despite the electrolytic liquid replacement. Slightly hyponatremic findings were accompanied with edema in the hands, face, and feet. As of the 7th day of observation, AST and ALT values turned back to normal rates, and the LDH levels could hardly return to normal values as of the 10th day. As of the 10th day of observation, the hyponatremia and edema recovered. Throughout the monitoring duration, hypotension progressing in the case was not accompanied with any other hemodynamic disorders and findings of EPS. Furthermore, no respiratory distress occurred.

The case was externed on the 7th day of observation on the condition that she would be called for controls.

Discussion

Although the side effect profile and toxidrome of atypical antipsychotic drugs in adults are well known, their overdosage information in pediatric age groups is rather new and limited.

Recently, atypical antipsychotic agents have largely replaced traditional agents as first-line drugs for the treatment of schizophrenia and psychotic mood disorders. Considering the increase in atypical antipsychotics prescriptions and the increased risk of suicide in this patient population, the number of reported cases of antipsychotic drugs may be expected to increase. Olanzapine has been considered to be similar to clozapine, but olanzapine intoxication appears to have a relatively benign clinical course as compared with clozapine intoxication. In olanzapine intoxications, deep coma, myosis, and
mild cardiovascular effects were observed (6).

According to Newcomer, the results of studies regarding second-generation (atypical) antipsychotics and metabolic effects are relevant to primary and secondary prevention efforts that aim to address the multiple factors that contribute to the increased prevalence of type 2 diabetes mellitus and cardiovascular disease in populations that are often treated with second-generation antipsychotic medications (7). Komossa et al. pointed out that olanzapine may be a somewhat more efficacious drug than some other second-generation antipsychotic drugs. This small superiority in efficacy needs to be weighed against a larger weight gain and associated metabolic problems than most other second-generation antipsychotic drugs, except clozapine. They suggested that further large, well-designed trials are necessary to establish the relative effects of different second-generation antipsychotic drugs.

Intoxication cases related with olanzapine are expected to increase day by day where olanzapine becomes a commonly prescribed antipsychotic medicine. In home child accidents, among 1-4-year-old children, intoxication with cleaning materials and drugs left around are encountered more frequently. The intake of olanzapine in the age range of 0-6 only occurs accidentally (5). The age and the way of intoxication of our case are in compliance with this fact. Especially in pediatric cases, the clinical experience in overdose of olanzapine and other atypical antipsychotics is limited. Research has demonstrated that the toxic dose for olanzapine is 0.5 mg/kg (enough to cause central nervous system depression). Unintentional antipsychotic ingestion in children can cause severe effects that last 1-3 days, often with one tablet.

Olanzapine's multireceptor action (antagonism to dopaminergic D1, D2, D4; serotoninergic 5-HT2A, 5-HT2C; histaminergic H1; cholinergic M1-5; and alpha-adrenergic receptors) results in multiple clinical symptoms in the course of acute poisoning. In the course of acute olanzapine poisoning, the prevailing symptoms come from the circulatory and central nervous systems; some symptoms are mutually opposed, such as coma-psychomotor agitation, hypertension-hypotension, tachycardia-bradycardia, hyperthermia-hypothermia, and miosis-mydriasis. Rarely, consciousness disturbances may persist for up to 6 days after olanzapine overdose; the course of poisoning can be severe, sometimes complicated, but fatal outcomes are rare. Kočer et al. (8) reported an adult case that developed diabetes insipidus following massive olanzapine ingestion and returned to normal after desmopressin treatment.

Overdose results in significant symptoms, including respiratory distress and mental status changes in infants. Catalano et al. (5) presented the case of an 18-month-old boy who ingested 30-40 mg of olanzapine, which resulted in significant symptoms, including respiratory distress and mental status changes. They stressed that previously reported pediatric cases of olanzapine ingestion have described similar symptoms. According to Catalano et al, the pediatric population should be monitored closely when ingestions of olanzapine occur. In another study, Catalano et al. suggested that special care may be warranted with all pediatric cases of atypical antipsychotic overdose, including monitoring in a high-intensity clinical setting to manage potential respiratory or cardiac difficulties.

Steil reported a case of delirium due to olanzapine overdose. After ingestion of 280 mg of olanzapine, a 19-year-old schizophrenic patient developed delirium with consciousness disturbance; disorientation in time, space, and situation; acoustic and visual hallucinations; and agitation. In this case, there were no abnormalities in the ECG, EEG, or routine blood tests. Approximately 36 h after the intoxication, the patient recovered fully. Furthermore, Steil pointed out that until now, there have been no reports of delirium from this cause. Theisen et al. reported on two cases of adolescents who attempted suicide with an overdose of olanzapine. A 14-year-old female ingested 275 mg olanzapine that caused somnolence, agitation (acutely), and EPS (after 54 hours) but no major clinical complications. A 17-year-old male ingested 400 mg olanzapine, which produced respiratory suppression requiring intubation and mechanical ventilation; he recovered after 3 days. Based on clinical monitoring and postmortem data, the two patients survived the ingestion of high doses of olanzapine. Morgan et al., in their study, described the spectrum of clinical effects in olanzapine overdose and investigated the factors that predict severe outcomes in adult patients. They analyzed olanzapine overdose events confirmed by drug analysis. In this study, demographic, clinical, and outcome
data were recorded for each presentation, and olanzapine overdose caused a high rate of delirium and central nervous system sedation that required significant resources. Morgan et al. suggested that olanzapine overdoses should be initially observed for 6 h; on the other hand, patients not taking olanzapine regularly may have more severe effects.

The initial findings of our case, such as lethargy emerging after the digestion of drug, may be based on the opiate-like effect of olanzapine. The hypotension and tachycardia observed in our case seem to be based on the α-adrenergic receptor occupancy property of olanzapine. In these cases, serum olanzapine level, weight of the patient, coingestants, the health of the patient at baseline, relevant laboratory and toxicology studies, and a standardized scale to rate the level of consciousness, such as the Glasgow Coma Scale (GCS) and the Poisoning Severity Score (PSS), should be assessed. On admission, in our case, GCS was slightly low (GKS=12). Furthermore, clinically, agitation and drowsiness were seen somewhat.

The most frequently employed treatments include intubation, gastric lavage, activated charcoal, intravenous fluids, artificial respiration, and restraints or sedatives. It is a more convenient approach to apply gastric lavage up to 2 hours later and generate absorbents effect by activated charcoal up to 6-8 hours later than digestion of medicine in cases of intoxication of medicine. In our case, we thought that applying gastric lavage 7 hours later after digestion would still be effective with respect to the probable anticholinergic effect of olanzapine. As the hypotension did not cause hemodynamic disorder in our case, crystalloid liquid replacement was decided to be sufficient, as sympathomimetic agent usage was not needed. However, throughout the observation, temporary rhythm disorders and T wave alterations that emerged in ECG of the case might imply arrhythmogenic and cardiotoxic effects of olanzapine. Tan et al. found 13 pediatric cases (<7 yr), 22 adolescent cases (7-16 years), and 185 adult cases. No pediatric case described a ventricular dysrhythmia or a cardiovascular death. In the adolescent and adult cases, we found numerous reports of prolonged QTc interval and hypotension, but there were only three cases of ventricular dysrhythmia and three deaths that may have been due, to direct cardiovascular toxicity. The results from case series reports were similar to the single case report data. Their findings suggested that overdose of atypical antipsychotics is unlikely to cause significant cardiovascular toxicity. However, further research about this subject is needed. Heart rhythm alterations showed the necessity of close cardiovascular observation in cases as well.

Ballesteros et al., in their study, reported an acute olanzapine mono-intoxication with severe toxicity and high whole-blood olanzapine concentrations. They pointed out that clinical and analytical data of similar samples obtained in non-fatal life-threatening cases can be very useful when interpreting postmortem cases.

Olanzapine is an atypical antipsychotic with multireceptor affinity and different pharmacological effects, which can result in abnormalities in laboratory investigations. In acute olanzapine poisoning, muscle and liver injury, serum glucose and electrolyte abnormalities, and changes in complete blood count (CBC) can be present; the valuable parameters for the monitoring of the course of poisonings are: serum activity of creatine phosphokinase (CPK) and transaminases (alanine transaminase: AST, alanine transaminase: ALT), serum level of bilirubin, glucose, potassium, and sodium and CBC. Hyperprolactinemia probably lacks practical importance, but further investigations are needed in this area. Waring et al. reviewed case notes from 64 consecutive patients admitted to their own institution after olanzapine overdose. They demonstrated overall and serum CK that was higher than 5 times the upper limit of normal in 17% of patients. According to this study, the prevalence of raised creatine kinase (CK) values was positively correlated with the stated quantity of olanzapine ingested, suggesting a dose-dependent relationship for acute muscle toxicity, and despite the high prevalence of acute muscle toxicity after olanzapine ingestion, none of the patients developed renal failure. In our case, serum CK levels were normal. No acute muscle toxicity after olanzapine ingestion was found. This state may be related to lower-dose olanzapine ingestion.

In our case, the high levels of AST and ALT determined 7 hours later then digestion of drug were related with conjugation and oxidation of olanzapine in liver (1). It might indicate that the elimination half-life of olanzapine in babies is a bit longer, and the lactate dehydrogenase (LDH) enzyme plays a part in the metabolism of medicine, as the AST and ALT levels could hardly return back to normal values after the 7th day and LDH levels could hardly return back to normal values after the 10th day of observation. The accomplishment of slight hyponatremia to edema beginning on the 3rd day of observation and the simultaneous recovery of both diseases might imply that hyponatremia is an edema-causing side effect of olanzapine. Further research about this subject is needed. Throughout the monitoring duration, the accompanying of hemodynamic disorders with hypotension, findings of EPS, or respiratory distress did not occur, as we interpreted them to be relevant to lower dose intake of olanzapine. Except for the extreme overdose cases, emerging clinical findings are considerably reversible; yet, close monitoring is recommended. Rare fatal pediatric cases have been recorded, as well (3,10). Lennestal et al. reported that in their study’s series of serum concentrations, a two-phase elimination was seen, with an initial elimination half-life of about 24 h during the first 3 days, followed by a second phase with a half-life of about 2.5 days. The patient in this case recovered completely. Because the elimination time after intoxication can be considerably longer than expected, it is recommended that the patient’s serum concentrations after intoxication be monitored.

In a multicenter study, the annual reports of the American Association of Poison Control Centers National Data Collection System were reviewed from 1990 to 2003, the most recent report currently available. All fatalities in children and youths under 18 years of age were included. In this study, the literature review identified 40 reports that included 63 patients, ranging in age from 1 day to 17 years of age. The clinical presentations included drowsiness,
lethargy, agitation, irritability, combativeness, and tachycardia. There were 11 fatalities in the cases reviewed: 1 from clozapine overdose, 3 from risperidone overdose, 2 from olanzapine overdose, and 5 from quetiapine overdose. All other cases reported no significant sequelae and resolved without any reported clinical consequences. Duration of overdose symptoms ranged from 24 hours to 7 days. One case of clozapine intoxication showed resolution of symptoms in 6 hours, and in another case of olanzapine overdose, symptoms resolved in 13 days. This study suggested that it is a need for future case reports to include serum medication level, weight of the patient, coingestants, the health of the patient at baseline, relevant laboratory and toxicology studies, and a standardized scale to rate the level of consciousness, such as the GCC. On the other hand, the existing pharmacovigilance data reports indicate that these medications are relatively safe when taken in overdose, particularly when coingestants are not involved.

Cobaugh et al. prepared a guideline to assist poison center personnel in the appropriate out-of-hospital triage and out-of-hospital management of patients with suspected acute ingestions of atypical antipsychotic medications by describing the process by which an ingestion of an atypical antipsychotic medication might be evaluated, identifying the key decision elements in managing cases of atypical antipsychotic medication ingestion, providing clear and practical recommendations that reflect the current state of knowledge, and identifying needs for research.

Balicka et al. presented a retrospective analysis of the clinical course of eight acute olanzapine intoxications treated at the Department of Clinical Toxicology, Jagiellonian University Medical College. In this study, central nervous system (CNS) symptoms, manifested as fluctuations between somnolence/coma and agitation/aggression and miosis, were observed in most of the patients. Moreover, increased CPK activity was stated in the most of patients. Consequently, all of the patients recovered, and poisoning severity according to PSS was moderate and severe (10). In our case, PSS was moderate (PSS=2). This state may be related to lower-dose olanzapine ingestion.

Chue and Singer examined the available data on olanzapine in untreated overdose situations. They point out that olanzapine is associated with toxicity in certain overdose situations, but evidence of any relation is limited and likely influenced by the higher rates of cardiovascular disease and sudden death in subjects with schizophrenia. This study suggested that similar toxicity data reviews should be conducted for all commonly prescribed psychotropics. In addition, early signal detection and effective notification processes are crucial in the event that serious adverse effects do occur.

Komossa et al. pointed out olanzapine may be a somewhat more efficacious drug than some other second-generation antipsychotic drugs. This small superiority in efficacy needs to be weighed against a larger weight gain and associated metabolic problems than most other second-generation antipsychotic drugs, except clozapine. They also suggested that further large, well-designed trials are necessary to establish the relative effects of different second-generation antipsychotic drugs.

Although olanzapine is tolerated relatively well in acute overdose, unpredictable and transient fluctuations between central nervous system depression and agitation, frequently associated with miosis, appear to be characteristic findings in moderate to high olanzapine overdoses. They are transient in nature and require careful clinical monitoring but rarely require specific therapeutic interventions.

**Conclusion**

Having side effects on the central nervous system, cardiovascular system, and metabolism, olanzapine can threaten lives, depending on dosage, especially accidental ingestion cases in children. Parents should consider that not only olanzapine but also all other medications left around means danger for children. Therefore, it is important to inform families sufficiently about this subject with respect to preventive medicine. This case is important in terms of olanzapine to take its cardiac effects in pediatric cases, even in low-dosage digestions. Olanzapine seems to be a medium-level safe medicine in terms of overdosage.

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**References**


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