Convulsive syncope related to a small dose of quetiapine in an adolescent with bipolar disorder

Abstract: Quetiapine, an atypical antipsychotic, has been extensively used in patients with bipolar disorder. Overdose of quetiapine can result in severe complications, such as coma, seizure, respiratory depression, arrhythmia, and even death. However, the paucity of toxicological evaluation in adolescence causes more potential risks in this population. Herein, we present a case of hypotension and convulsive syncope after exposure to a small dose of quetiapine in a 16-year-old who was diagnosed with bipolar disorder. After cessation of quetiapine, no additional convulsive movements were reported. This case indicates that even in young patients without predisposing factors, close monitoring of adverse effects should be warranted for safety concerns, especially at the initiation of quetiapine treatment.

Keywords: quetiapine, bipolar disorder, hypotension, convulsive syncope

Introduction
Quetiapine, a second-generation antipsychotic, is approved by US Food and Drug Administration to treat young patients (age ≥10 years) with acute depressive or manic episodes and maintenance of bipolar disorder.¹ Quetiapine seems to be a promising agent, well tolerated in the short and medium term in children and adolescents, and has low potential to elicit extrapyramidal effects.² However, extremely limited guidelines for its use are available in this population. A recent nationwide report demonstrated that off-label use of quetiapine in children and adolescents is prevalent in Denmark, and life-threatening adverse events involved seizure, QT-interval prolongation, and cerebral hemorrhage.³ Besides, this study implied that quetiapine could induce hyperprolactinemia, especially in minors.³ Moreover, in some susceptible individuals, exposure to quetiapine overdoses can result in severe complications, such as coma, seizure, respiratory depression, arrhythmia, and even death.⁴ Therefore, great care should be taken when initiating quetiapine treatment in young patients.

For example, new-onset seizure has been reported in two young patients after taking small doses of quetiapine.⁵ One of these two patients was a 16-year-old girl with ornithine transcarbamylase deficiency and mental retardation, who was prescribed quetiapine to manage her behavioral instability. She developed a generalized tonic–clonic seizure after ingesting 50 mg quetiapine for 2 consecutive days.⁶ Another was a 7-year-old boy with autism and mental retardation, who was prescribed quetiapine at a dose of 25 mg/day for his behavioral problems. Generalized tonic–clonic seizure occurred as early as the first time quetiapine was taken.⁶ According to the literature available, most cases of seizure secondary to quetiapine exposure occur in adult patients or present as acute intoxication because of overdoses.⁶–⁸ Evidence of severe adverse events associated with quetiapine treatment in children and adolescents is fairly rare.
Herein, we depict an adolescent with bipolar disorder who developed hypotension and syncope with convulsive movements after ingestion of quetiapine without concomitant medications. This case enriches our knowledge of quetiapine toxicity in young people.

Case presentation

A 16-year-old high school teenager was admitted to our hospital because of episodic mood swings for almost 4 years. Periods of depression were predominant in his course of disease and lasted for 1 or 2 months. When depressed, this patient complained of waning interest, poor academic performance, bad memory, and loss of appetite. Sometimes, this patient would have suicidal ideations, and had once attempted suicide by holding his breath. He had auditory hallucinations and delusions of persecution. Comparatively, when his mood elevated, this patient would become energetic, conversant, and generous, and gradually became euthymic after nearly 1 week. Two weeks before admission, this patient was sent to the outpatient service by his family. No physical illness, epilepsy, history of head trauma, or family history of mental disorders was reported. He had lost his mother because of a traffic accident. He denied experience of smoking, drinking alcohol, or taking any toxic or illicit substance. Neurological examination was normal, as well as laboratory tests (e.g., routine blood test, urine, thyroid hormones, reproductive hormones, biochemical indices, and infectious profiles). Magnetic resonance imaging of the brain indicated an enlarged left lateral ventricle (Figure 1). According to the Diagnostic and Statistical Manual of Mental Disorders (fifth edition), this patient matched with the diagnosis of bipolar II disorder and depressive episodes, with psychotic features. Consequently, quetiapine was prescribed and started at a dosage of 50 mg per night. However, when this patient took the first 50 mg, he slept for almost 12 hours and felt feeble in the daytime. Since then, he had decided to stop this medication. His depressive condition did not improve, and hospitalization was suggested by his psychiatrist.

On admission, all vital signs of this patient were normal. His grandmother stayed in hospital to look after him. This patient tried to take an afternoon nap, but failed to fall asleep, and was given 50 mg quetiapine as a sleeping aid by his grandmother. About 2 hours after quetiapine exposure, this patient was taken to the nurses’ desk and educated with notes on security as a new inpatient. All of a sudden, his complexion became pale. Immediate blood glucose was 5.5 mmol/L, heart rate was 72 beats per minute, and blood pressure 118/80 mmHg. The psychiatrist arrived quickly and tried to measure blood pressure again, but failed to get any result. On his way back to the ward when supported by others, this patient fell abruptly to the floor, followed by tics of limbs and opisthotonos, which lasted for 2–3 seconds, and lay on the floor unconscious. No foaming at the mouth, urinary incontinence, or tongue biting was observed. About 1 minute later, he stood up by himself and asked, “What happened just now? My mind went blank.” At this moment, his complexion gradually became fresh and rosy. Blood pressure was at 88/53 mmHg, and heart rate was at 61 beats per minute. As the patient’s condition had improved, he was given low-flow oxygen therapy with close electrocardiography monitoring. Serum creatinine kinase after this event was 1,960 units/L (reference range 38–174 units/L). We did not examine the quetiapine concentration in the blood, as the patient had taken a small dosage of this medication. He was told not to take this medication again. A combination of valproate and olanzapine was used as his therapeutic regimen. Electroencephalography (EEG) was conducted 5 days after the event, and no epileptiform activity was captured. This patient recovered well in the next 2 weeks and remained syncope-free before hospital discharge. During outpatient visits, no more convulsive syncope was reported.

The Institute Ethical Committee of the First Affiliated Hospital, Zhejiang University School of Medicine approved this case study. Written informed consent for publication of the case and any accompanying image was obtained from the patient and his guardian.

Figure 1 Magnetic resonance imaging showing an enlarged left lateral ventricle (orange arrow).
Discussion

In this study, we present a case of extreme hypotension, transient convulsive movements, and unconsciousness after quetiapine exposure, suggesting relevant adverse effects to a small dosage of quetiapine (50 mg) in an adolescent with bipolar disorder. No predisposing factors were identified in this patient. To our knowledge, this is the first case study suggesting convulsive syncope secondary to a small dosage of quetiapine (50 mg) in an adolescent with bipolar disorder.

Quetiapine functions as a multireceptor antipsychotic involved in the dopamine, serotonin, and adrenergic pathways. It also has antihistamine and anticholinergic properties. Hypotension is known as a common adverse effect, which is considered to be elicited by inhibition of adrenergic α1-receptors. However, the mechanism behind quetiapine-induced seizure remains unclear. Structurally similar to quetiapine, the dose-dependent potential of clozapine to mediate seizure by lowering the threshold of seizure is more definite. The risk of seizure provocation induced by first-generation antipsychotics or second-generation antipsychotics is inconsistent across different studies. In a nest-control study, when applied to patients with affective disorders, medium- to high-potency first-generation antipsychotics were more likely to elicit seizure than all other antipsychotics. Nevertheless, other findings suggested second-generation antipsychotics, rather than first-generation antipsychotics, had a higher risk of seizure. In addition, abnormal alterations on EEG have been observed in psychiatric patients taking second-generation antipsychotics, which were particularly high under clozapine and olanzapine treatment. None of the patients with quetiapine treatment showed EEG abnormalities in the same study. Interestingly, in a rat model with malformations of cortical development, quetiapine was found to rescue its cognitive impairment and reduce seizure susceptibility, possibly by maintaining the integrity of myelin.

In our patient, we noted that extremely low blood pressure was associated with convulsive movements. In individuals with quetiapine abuse, severe complications, such as hypotension, respiratory depression, and seizure, have also been observed. Unfortunately, the interrelationship between hypotension and seizure has always been neglected. In a rare case of seizure presenting as refractory hypotension, the authors provided sufficient evidence to support acute hypotension as an atypical manifestation of seizure. Hypotension during the onset of seizure may cause cerebral hypoperfusion and further worsen autonomic dysfunction.

Of note, we would like to differentiate the manifestations of convulsive syncope from those of seizure attacks. In our patient, acute onset of limb spasm, opisthotonos, and transient unconsciousness was observed without other typical symptoms of seizure, such as foaming at the mouth, loss of urine, or biting of the tongue. Therefore, the seizure-like movements in our patient were not typical, and were more likely to be a syncopal attack secondary to hypotension. Convulsive syncope, or in academic terms, reflex asystolic syncope, occurs predominantly in young people and can manifest as loss of consciousness, loss of postural tone, myoclonic movements, and nonepileptic spasms.

In the present case study of a patient with bipolar disorder, hypotension associated with convulsive movements was observed nearly 2 hours after exposure to a small dose of quetiapine. This is in accordance with the serum peak time after taking quetiapine orally. No obvious predisposing factors were reported, and no other medications were congested. However, the undetected quetiapine concentration in blood seemed to be a major limitation of our study. In individuals with specific polymorphisms in CYP3A4 enzymes, unpredictable adverse reactions were associated with decreased enzyme activity and increased blood concentration. Moreover, no additional convulsive movements were observed after discontinuing quetiapine treatment. According to a scale evaluating possible drug-related adverse effects, the relationship between reported adverse reactions and quetiapine in our patient was assigned to the “probable” category. Finally, we hypothesize that the convulsive movements were consequences of hypotension induced by quetiapine.

In conclusion, this single case report enriches our understanding of quetiapine-associated adverse reactions in young patients. Special concerns are essential to evaluate the risk of hypotension and convulsive syncope in children and adolescents before starting quetiapine treatment. Continuous monitoring of acute adverse events is also needed after the initiation of treatment.

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