Restless legs syndrome following the use of ziprasidone: a case report

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ABSTRACT
Restless legs syndrome (RLS) is a common sleep-related movement disorder characterised by an uncomfortable urge to move the legs that occurs during periods of inactivity. Although there have been many case reports on antipsychotic-induced RLS, ziprasidone has never been reported as a cause of RLS. We present a case of a female patient with schizophrenia who presented with symptoms of RLS following the administration of high doses of ziprasidone added to quetiapine and valproate. The patient's symptoms of RLS occurred following the administration and titration of ziprasidone to 160 mg, and were relieved upon reducing the dose to 120 mg/day. Other potential causative medications and differential diagnoses that could have caused similar symptoms were excluded. Clinicians should be aware of the potential for ziprasidone-induced RLS. Dopamine and serotonin interaction could be the mechanism underlying ziprasidone-induced RLS.

INTRODUCTION
Restless legs syndrome (RLS), with prevalence ranging from 2% to 15% in the general population, is a common disorder associated with many medical conditions and medications. Some second-generation antipsychotics such as risperidone,1 olanzapine2 and quetiapine3,4 have been reported to cause secondary RLS, whereas others have been reported to help relieve it.1,3 However, no cases of RLS induced by ziprasidone were found on MEDLINE or EMBASE searches done on 25 March 2018. We report a case of RLS possibly induced by ziprasidone.

CASE HISTORY
Introduction of the case
Miss A was a 35-year-old Chinese female patient with a history of known mild intellectual disability since childhood and an 8-month history of schizophrenia (Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition). She was initially treated with risperidone titrated to 6 mg orally per day (3 mg two times per day) with partial improvement but persistent active commentary auditory hallucination, disorganised behaviour and agitation. A decision was made to switch to quetiapine with a maximum dose of 800 mg daily (400 mg orally two times per day) and 500 mg of valproate acid daily. While taking 800 mg of quetiapine daily, the patient still reported residual symptoms of paranoia and intermittent auditory hallucination, so we decided to put her on ziprasidone. We started with 40 mg daily and increased to 40 mg every 3 days (40 mg–80 mg–120 mg–160 mg). She developed significant improvement of psychotopic symptoms after the addition of ziprasidone. However, the night after she was titrated to 160 mg daily (80 mg at lunch and 80 mg at dinner) of ziprasidone, she developed marked, intractable discomfort in bilateral legs, which she could not describe clearly given her poor iteration. The discomfort induced an urge to move her legs, was partially relieved by walking, and worsened with resting. The symptoms were worse at night and in the afternoon. She was unable to fall asleep due to significant discomfort and kept pacing in the unit during the night. Benzodiazepines were prescribed as needed to help relieve her symptoms.

Diagnosis of RLS
A diagnosis of RLS had been made based on the diagnostic criteria from the International Restless Legs Syndrome Study Group (IRLSSG).6 Miss A’s symptoms met all the criteria of this IRLSSG guideline (5/5). As for the severity of symptoms, the patient rated her symptoms as Severe (30 points in total) using the International Restless Legs Syndrome Study Group Rating Scale (IRLS). We also ruled out any diagnosis that could mimic symptoms of RLS, making sure that the preceding manifestations were not solely accounted for as symptoms primary to another medical or behavioural condition such as myalgia, venous stasis, leg oedema, arthritis, leg cramps, positional discomfort or habitual foot tapping.

Akathisia was ruled out on the basis of the presence of dysesthesias in the legs rather
than gross restlessness, and worsening symptoms at night. The neurological examination (cranial nerves, motor system, reflexes, sensation, and so on) of the patient was unremarkable, so movement disorders could be ruled out. Anaemia (complete blood count), iron deficiency (complete iron panel), diabetic neuropathy (fasting blood glucose), vitamin B₁₂, and folate deficiency had been ruled out. Renal function (blood urea nitrogen, creatinine), thyroid function (thyroid-stimulating hormone, FT₃, FT₄, T₃, T₄), liver function (aspartate aminotransferase, alanine aminotransferase, bilirubin, direct bilirubin, gamma-glutamyl transferase) and brain MRI were all within normal limits. Per patient and collateral, Miss A has no history of smoking, alcohol or substance use, nor history of excessive caffeine use. History or family history of similar manifestations or movement disorders was also ruled out per patient and collateral history.

Management and clinical course after the onset of RLS
As several reports have shown that RLS could be induced by quetiapine, we tapered Miss A off quetiapine and valproic acid. However, even with ziprasidone as monotherapy, her symptoms of RLS persisted.

Given the patient’s amelioration of psychotic symptoms with ziprasidone, treatment with ziprasidone was maintained but with a reduced dose of 120 mg daily (40 mg three times a day during meals). The day after we reduced the dosage, the patient reported minimal discomfort in bilateral legs in the afternoon, which was relieved with massage within an hour. She reported sufficient sleep at night after tapering down the dose of ziprasidone. The patient rated her symptoms as Mild (9 points in total) at this time using the IRLS. Several days later, Miss A’s psychotic symptoms were resolved, and she was thus discharged home.

DISCUSSION
The presentations of our case met the diagnostic criteria established by IRLSSG for RLS. We differentiated RLS from akathisia by its manifestations. However, RLS is often associated with periodic limb movement disorder (PLMD) that can only be diagnosed with polysomnography. Unfortunately, polysomnography was not able to be done due to resource limitations in the hospital, thus the association with PLMD was unable to be ascertained. The patient developed RLS on ziprasidone dosing 160 mg daily and symptoms were significantly relieved on reduction to 120 mg daily (optimal effective dose), divided three times a day. Our patient developed short duration of RLS daily but was able to tolerate it.

Since ziprasidone is a minor substrate of cytochrome P450 (CYP)3A4 and CYP1A2, and quetiapine is metabolised primarily by CYP3A4 and valproic acid by mitochondrial beta-oxidation, the drug interactions between them should be minimal.

To the best of our knowledge, considering the growing use of ziprasidone worldwide for the last two decades, no case had been reported regarding RLS and ziprasidone to date.

Dopaminergic dysfunction is considered to be a possible pathophysiological mechanism in the development of RLS, and drugs that antagonise dopamine receptors are likely to induce RLS. Antipsychotics such as ziprasidone generally have intermediate to high D₂ dopamine receptor occupancy. However, within an environment of 5-hydroxytryptamine (5-HT) elevation, it was likely 5-HT₁A hyperactivation or 5-HT₂A antagonism, which potentiated the serotonin-dopamine imbalance, contributed to dopaminergic hypactivity and RLS development in our patient. What’s more, like other atypical antipsychotic agents, ziprasidone is a 5-HT₂A/dopamine D₂ antagonist. However, its in vitro 5-HT₂A/D₂ receptor affinity ratio is higher than that of the other first-line atypical antipsychotic agents (namely risperidone, olanzapine, quetiapine and aripiprazole). Moreover, ziprasidone has relatively high affinity on 5-HT₁A receptor, acting as a partial agonist. Paroxetine worsened cases of RLS as in previous reports. It is speculated that an increased serotonergic activity leading to an indirect dopaminergic antagonism by paroxetine could be the explanation of RLS generation.

It is interesting to note that as a potent dopamine blocking agent, risperidone did not induce RLS in our patient. A possible explanation could be that our patient was a fast metaboliser of risperidone, which is metabolised by CYP2D6. Another explanation might be due to the alterations in the serotonergic neurotransmission system alone, for serotonin plays a critical role in motor function and sensory information processing. If that is the case, we can assume that a more potent 5-HT₁A activating agents, like cariprazine, adoprazine, could be more likely to aggravate RLS symptoms than a less potent one.

In conclusion, we found a dose-related case of RLS in a patient on ziprasidone therapy. The pathophysiology remained obscure. Further studies are needed to explore the mechanism of dopamine dysfunction underlying psychotropic-induced RLS. Additionally, clinicians should be aware of the possibility that ziprasidone could develop RLS, and that the severity of RLS symptoms could be dose related. Discontinuation or tapering down the dosage may be needed to relieve the symptoms of secondary RLS.

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Reference
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