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A case report

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Clozapine was the first atypical neuroleptic to be developed, and has been used since the 1960s. It is well known as the treatment of ‘last resort’.1,2 In clinical practice there is often reluctance to switch individuals to this medication because of its side-effect profile, despite its proven efficacy even in treatment-resistant schizophrenia.3

The adolescent and child psychiatry unit at the University of Malaya Medical Centre, Kuala Lumpur, has a section that specifically caters for children diagnosed with early-onset psychosis. The unit is currently treating 3 adolescents with clozapine (the number is increasing). One of these cases will be described in this report.

Case report
A 15-year-old Chinese girl was brought to the unit by her parents in 2008. She had been treated by a private psychiatrist for over a year, and had been receiving an atypical antipsychotic. Her parents felt that she had done fairly well initially, but her condition worsened. She was suspicious of her classmates and felt that they were constantly making comments about her, and also complained that she could not trust her friends. Her parents were worried that she would act on these suspicions. She was irritable and had a lot to say when spoken to. She also complained that her eyelids were not wide enough and that she needed to go for an operation to widen them. She wore a lot of make-up and thick clothing. She also had sleep problems, staying up at night, reading magazines and watching television until the early hours of the morning.

During the mental state examination the patient denied experiencing any auditory hallucinations, but was noted to be speaking to herself.

Even with good compliance with medication the patient’s symptoms worsened. It was decided that the best option was to switch to another atypical antipsychotic. However, after 2½ months, although she was sleeping better, she became more aggressive and her parents were afraid of her frequent outbursts. It was decided to try another atypical antipsychotic, but despite increasing the dose of the medication and allowing sufficient time for it to have an effect, no improvement was noted. It was then decided to try clozapine.

The patient had had no previous medical illnesses of note. There was no history of substance use, and no significant family history of mental illness. A computed tomography scan of the brain and the results of relevant routine blood tests were normal.

After a week on clozapine, the patient seemed calmer and began to socialise appropriately with her family. She remained in the ward for another week and was eventually discharged on 150 mg clozapine at night. During subsequent follow-up she was found to be calm and had no psychotic features. She was eventually able to start going to school again. Her parents were happy with her improvement, and her blood parameters remained within normal limits.

Discussion
Early-onset schizophrenia is defined as onset before the age of 18 years, with very early onset before the age of 13 years.4 It is often a terrifying and debilitating condition, and costly for the child and the family. Reports have suggested that the outcome is not generally favourable and that it is a progressively deteriorating developmental disorder, with up to 25% having a full remission with a good prognosis, another 25% showing partial remission, and up to 50% becoming chronic.4-6 Individuals receiving treatment for a first episode of schizophrenia often demonstrate a lasting response to antipsychotic medications, though a subgroup are left with significant persisting psychotic symptoms.7-9 The focus during recent years has been on improving understanding of the illness and ultimately more effective intervention.

Reports have demonstrated the efficacy of clozapine.10,11 Unfortunately the therapeutic benefits may be accompanied by
severe adverse effects in children. The most serious side-effect is bone marrow suppression and agranulocytosis, which has delayed the acceptance of clozapine into clinical practice. However, failure to treat children who do not respond to routine treatment because of a fear of side-effects is not justifiable, as the risk of not treating schizophrenia is more serious than the adverse effects of treatment. Furthermore, a delay in treatment is often associated with poor results, as the longer the duration of untreated psychosis the worse the condition becomes, with functional outcome declining sharply. The persistent negative as well as positive symptoms are destructive to a child’s development, to progress at school and to family and peer relationships, as was demonstrated in our case. The suicide rate in patients with childhood-onset schizophrenia has also been noted to be higher than in non-schizophrenics. Early treatment therefore has the potential to reduce the secondary impacts of this serious mental illness. In our case the onset of frequent episodes of aggression, which significantly affected the child’s functioning, as well as the imminent potential harm to others indicated a need for immediate pharmacological intervention. Clearly the symptoms were not responding to the antipsychotics used.

Clozapine has value in the salvage of patients considered resistant to treatment, even in the paediatric population. It is the only drug licensed for the treatment of schizophrenia in individuals as young as 16 years who are unresponsive to or intolerant of conventional medication. Treatment-resistant patients are those who fail to respond to trials of two typical antipsychotics, or in this age of newer atypical antipsychotic agents who fail to experience a substantial reduction in psychopathology with at least three different antipsychotic agents.

Although clozapine has the potential to cause severe side-effects, there is increasing evidence to advocate its use in the early treatment of first-episode patients whose psychosis does not remit with other second-generation antipsychotics. It has been suggested that clozapine should be positioned as a second-line treatment for first-episode schizophrenics who fail one trial of a second-generation antipsychotic.

The most frequent side-effects observed are drowsiness, drooling, nonspecific excitatory changes on the electroencephalogram (EEG), transient psychomotor agitation and eosinophilia. General guidelines for using clozapine in children are the same as for adults, but since there is a higher rate of side-effects in children, clozapine must be used with extra caution in the paediatric population, and close monitoring for adverse events is essential. Monitoring of the white blood cell count is similar to that in adults, i.e. blood counts should be done weekly for the first 6 months of treatment and every second week thereafter. It is prudent to do a baseline test, and to include a baseline EEG. It is also recommended that lower doses of antipsychotics than in adults be used to achieve a therapeutic response. Start low, go slow, taper slow’ dosing strategies are especially important when atypical antipsychotic agents are tried on adolescents. The long-term outcome in treatment-resistant children seems to be predicted best by the improvement seen at 6 weeks, by which time most progress will have occurred.

Bipolar disorder typically presents with affective and at times psychotic symptoms. Mania in adolescents can therefore present with psychotic symptoms, i.e. delusions, hallucinations and thought disorder. Our patient demonstrated prominent delusions of reference and persecution, and although she denied hearing any voices, she was seen talking to herself. There were no overt manic or depressive symptoms, although she did display irritability. The diagnosis of schizophrenia rather than a psychotic mood disorder was therefore made. However, longitudinal assessment is needed to ensure accuracy of diagnosis in all such cases.

As our child mental health services increase, we are likely to see more children with early-onset treatment-resistant schizophrenia. It is hoped that this report will add to the current body of knowledge and encourage clinicians in this part of the world to use clozapine for the benefit of our child patients. More than 15 studies have demonstrated the antipsychotic efficacy of clozapine in childhood and adolescent schizophrenia. The main advantages of clozapine treatment in this age group in comparison with typical antipsychotics are: (i) good antipsychotic efficacy during an acute schizophrenic episode; (ii) better improvement in chronic cases with a high load of negative symptoms; and (iii) markedly fewer extrapyramidal adverse effects, and therefore fairly good tolerability.

References


