Closure beyond clozapine: successfully averting rebound symptoms in a patient with schizoaffective disorder and agranulocytosis

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Summary

‘Rebound’ or ‘withdrawal’ symptoms are frequently observed after a sudden discontinuation of clozapine. We describe a patient with treatment-resistant schizoaffective disorder who developed agranulocytosis on clozapine but was successfully switched to treatment with olanzapine with no deterioration in her condition. We put forward three possible theories which may have accounted for the lack of rebound symptoms in this patient: the pharmacological profile of olanzapine, the anticholinergic effects of hyoscine hydrobromide, and the possibility that this patient may not be treatment-resistant and so have a reduced risk of rebound psychosis due to displaying a different pathophysiology.

Declaration of interest

None.

Keywords

Clozapine; treatment-resistant schizophrenia; rebound psychosis; agranulocytosis; psychosis.

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Case presentation

A 72-year-old White woman, Ms G, with a 40-year history of treatment-resistant schizoaffective disorder had been managed as an in-patient for the past 6 months under Section 3 of the Mental Health Act. She has described prominent persecutory and grandiose delusional symptoms over the course of her illness, in addition to irritability, mood instability, formal thought disorder, self-neglect and lack of insight. Among Ms G’s prominent beliefs are that she is being tricked into paying for her National Health Service care, that she is the victim of identity theft and has several children, and that her psychiatric symptoms are the result of being spiked by drug dealers.

She had been admitted to hospital several times over the course of her illness, demonstrating a relatively consistent pattern of relapse, with prominent episodes of mania with psychotic features. Previously she had been treated with numerous oral and depot antipsychotic medications including trifluoperazine, haloperidol, aripiprazole, risperidone, olanzapine, pipotiazine, paliperidone and zuclopenthixol. A refusal to adhere with oral medications has been a characteristic feature of her illness in the past.

She experienced moderate to severe akathisia and extrapyramidal side-effects, particularly on the left side, on her depot pipotiazine palmitate, which was managed with procyclidine. Previously, she has had evidence of subclinical hypothyroidism.

Ms G has, during her adult life, had intermittent periods of heavy cigarette smoking, although had stopped since this admission to hospital. There is no recent history of drug use or harmful use of alcohol. The available records suggest that Ms G is single and has no children. Her mother died by suicide when she was 1 year old. Her father also died when she was a child.

Ms G was admitted to the National Psychiatry Service for a trial of clozapine because of poor symptom control when treated with other antipsychotics, and worsening extrapyramidal symptoms on depot antipsychotics. On admission she was on a weekly depot of zuclopenthixol (500 mg). This was discontinued, and clozapine was titrated up to 250 mg once daily. She responded well initially. An improvement in her mood and a reduction in her irritability and delusional beliefs were noted. However, in her eighth week of treatment her neutrophil levels began to fall, reaching 0.38 × 10^9/L (white cell count 2.03 × 10^9/L), at which point clozapine was stopped, and the patient remained neutropenic until the fourth day after stopping clozapine, during which time she was hospitalised for the first time since admission. She experienced moderate to severe akathisia and extrapyramidal side-effects, particularly on the left side, on her depot pipotiazine palmitate, which was managed with procyclidine. During this time, she was clinically well with no signs of infection. Other medications...
taken at the time were hyoscine hydrobromide (300 μg three times a day as required) and cholecalciferol (800 units every morning).

Olanzapine was then started (10 mg) at night on the seventh day after clozapine was stopped. There was no deterioration in her mental state during this time off antipsychotic medication. The rationale for this choice of medication was that she had previously responded well to olanzapine during a 3-month period of treatment with subsequent relapse, thought to be related to non-adherence, but not confirmed. At review after 2 weeks on the current trial of olanzapine, no delusions were elicited, there was no evidence of formal thought disorder and she reported a better clarity of mind and reduced hypersalivation than on clozapine. The dose was later increased to 12.5 mg after some increase in delusional beliefs were noted, but she has now been stable for 3 months and is being discharged to a rehabilitation unit.

Discussion

Averting clozapine rebound symptoms: olanzapine, anticholinergics or neurochemical profile?

The mechanism causing the withdrawal symptoms associated with a sudden discontinuation of clozapine is unclear. It has been postulated that they are due to clozapine-induced ‘supersensitivity’ of receptors for dopamine, acetylcholine, gamma-aminobutyric acid and serotonin. Thus, these symptoms occur owing to rebound increase in the corresponding neurotransmitter activity.6 Clozapine withdrawal has been suggested to result in psychotic relapse more than some other antipsychotics. Clozapine action via looser binding and rapid displacement at the D2 receptor by endogenous dopamine is another theory that has been posited to explain this rapid withdrawal symptoms.7 There is no robust data on incidence of clozapine withdrawal symptoms, but one retrospective study of 28 patients found that 46.4% of patients experienced rapid deterioration in mental state.8 It is therefore an important consideration when circumstances dictate a sudden halt to clozapine therapy. We put forward three possible theories that may have accounted for the lack of rebound symptoms in this patient.

Switching to another antipsychotic medication is a possible approach after abrupt clozapine cessation, although often unsuccessful. Treatment is often symptomatic, using measures such as anticholinergics for dyskinesia and dystonia,9 and electroconvulsive therapy for catatonia.10 In terms of antipsychotic medication, there is insufficient evidence to favour any particular drug. However, olanzapine in above-licensed doses is most commonly used.5 Two randomised controlled trials comparing olanzapine with clozapine in treatment-resistant schizophrenia found olanzapine to have similar efficacy.11,12 This and may be owing to its similar pharmacological profile to clozapine.13 However, the Maudsley Prescribing Guidelines note that clinical experience of substituting clozapine is disappointing.6 One study has specifically compared olanzapine with placebo in patients where clozapine has been discontinued.14 The placebo was only given for 3–5 days before commencing olanzapine, but more patients treated with placebo (24.5%) than patients treated with olanzapine (7.5%) experienced clozapine discontinuation symptoms. Given that the patient in question had a longer, 7-day period between stopping clozapine and starting olanzapine, owing to agranulocytosis, it is possible that she would have been at greater risk of developing rebound symptoms than the patients in the placebo arm of this trial. However, the pharmacological profile of olanzapine may have been an important factor in this case once she had started taking it.

A further reason that may have influenced the successful outcome in this patient is the fact she was taking the anticholinergic medication hyoscine hydrobromide for hypersalivation. One theory of clozapine withdrawal phenomena is that they may be partly due to cholinergic overdrive after sudden removal of the strongly anticholinergic effects of clozapine.6 This explains symptoms such as nausea, vomiting, diarrhoea, headache, restlessness, agitation and sweating. In addition to somatic symptoms, anticholinergic withdrawal is also associated with an increase in pschotic symptoms. Patients receiving anticholinergic drugs have been found to be significantly less likely to deteriorate than those not receiving them.9 Consequently, anticholinergics have been recommended to treat relevant withdrawal symptoms both prophylactically and symptomatically.10 The other consideration in this patient is the fact that before clozapine she was on a depot zuclopenthixol. This has a half-life of 21 days and takes around 3 months to be eliminated. Hence, when the clozapine was stopped, she could still have had residual low levels of zuclopenthixol in her system.

It is also possible that this patient may not be treatment-resistant and so have a reduced risk of rebound psychosis owing to a different pathophysiology. It has been proposed that different treatment responses may represent different subtypes of schizophrenia.16 A recent meta-analysis17 found tentative evidence supporting the hypothesis that treatment resistance may be a separate disorder. In contrast to the dysregulation of the dopamine system in treatment-responsive schizophrenia, resistance was found to be better characterised by abnormalities in the glutamate system and reduction in grey matter. One potential explanation being that this patient may have responded well to olanzapine in the past, but only stopped complying covertly and relapsed because of non-adherence, meaning she may not have been genuinely treatment-resistant in the first place. Whether patients experience withdrawal symptoms after clozapine cessation may be a marker of a distinct neurochemical profile to their condition. However, she had been trialled with depot antipsychotic medication for a sustained period of time and still remained symptomatic, suggesting that this is not the entire story.

Of course, we cannot know for certain how or if these possibilities may have interplayed to produce a successful outcome in this case. However, they do raise important points in the management of clozapine withdrawal. More evidence is needed on the choice of antipsychotic and use of anticholinergics. Furthermore, distinguishing between non-adherence and treatment resistance in patients may be a predictor of who is likely to experience rebound symptoms. Current trials of digital medicine interventions may be helpful in this regard.18

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Consent statement

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