Is Clozapine to Blame for Delayed Ogilvie Syndrome and Gastrointestinal Bleeding in Overdose Settings?

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SUMMARY

Introduction: Ogilvie syndrome and gastrointestinal bleeding as complications after reversal of typical clinical picture of acute clozapine overdose is described.

Case Report: A previously healthy 31-year-old man was found unconscious with Glasgow Coma Score of 6, non-reactive miotic pupils, hypersalivation and heart rate of 115 bpm. In the blood, the presence of clozapine, diazepam, haloperidol and biperiden were confirmed. The patient was referred to the intensive care unit for symptomatic and supportive treatment. Clinical signs registered on admission, except for sinus tachycardia, were completely resolved by the day 3. The patient began to eat and had regular bowel movements. From the seventh day, gastrointestinal complications were noticed. Nonobstructive dilatation of the stomach and intestine was confirmed on computed tomography scan. Nasogastric suction, the usage of laxatives and prostigmin injections as well as colonic irrigation was performed with a good clinical response.

Conclusion: Clinicians should be aware of the potential of atypical antipsychotics to cause ileus, particularly in combination with other drugs with antimuscarinic properties, and ready to rapidly detect and treat intestinal atony thus preventing life-threatening complications. Serum clozapine levels may not equate to clinical toxicity and the drug-naïve patient require more careful observation for complications in clozapine toxicity settings.

Keywords: Atypical Antipsychotics, Clozapine, Acute Overdose, Delayed Complications, Ogilvie Syndrome, Gastrointestinal Bleeding

INTRODUCTION

Clozapine is an atypical antipsychotic with minimal extrapyramidal toxicity, indicated for the treatment of severely ill patients with schizophrenia who fail to respond adequately to standard acceptable antipsychotic medication [1]. However, its true potential in treatment-resistant schizophrenia is limited by the complex adverse effects’ profile [2].
Even though gastrointestinal hypomotility was described with typical antipsychotics mainly in association with the use of anticholinergic agents, some of the newer agents, such as clozapine, exhibit intrinsic anticholinergic activity as well [3]. Moreover, a recent study, by using a colonic transit test, provides objective evidence of significant gastrointestinal hypomotility in patients treated with clozapine [4].

Gastrointestinal hypomotility associated with clozapine may cause paralytic ileus, faecal impaction, aspiration of vomit, necrotizing colitis and/or intestinal perforation; fatalities provoked by those (often underrecognized) complications have been reported as well [5]. In addition, acute intestinal pseudo-obstruction (Ogilvie’s Syndrome), which is characterized by signs and symptoms of a mechanical obstruction of the small or large bowel in the absence of a mechanical cause, has been related to antipsychotics usage [6-8].

The authors point out that high toxic clozapine concentrations and subsequent paralytic ileus might be infection/inflammation induced in patients on chronic clozapine treatment or due to toxic clozapine ingestion (overdose) [9-15].

A case of a young man who developed gastrointestinal hypomotility and bleeding as complications after reversal of typical clinical picture of acute clozapine overdose is described here.

**CASE REPORT**

A previously healthy 31-year-old man was found unconscious with Glasgow Coma Score of 6, non-reactive miotic pupils, hypersalivation and heart rate of 115 bpm. Except for sinus tachycardia, the electrocardiogram was read as normal. Laboratory tests on admission were completely resolved by the day 3. The patient began to eat and had regular bowel movements.

At day 7, the patient complained of abdominal fullness and nausea, followed by vomiting copious quantities of tea-colored fluid. Physical examination showed distended abdomen and decreased bowel sounds. High leucocyte count (10.60x10^9/L – 24.0x10^9/L – 20.0x10^9/L; referent range: 4.0-10.0x10^9/L), with gradually falling levels of hemoglobin (100 – 93 – 80 g/L; referent range: 120-180 g/L) and hematocrit (0.31 – 0.27 – 0.24 L/L; referent range: 0.35-0.54 L/L) were detected. Under the suspicion of gastrointestinal perforation, a plain x ray of the abdomen was indicated. Findings, interpreted by an experienced radiologist, pointed to the existence of subdiaphragmatic free gas or the distended stomach. Nonobstructive dilatation of the stomach, duodenum, jejunum and the proximal parts of ileum was confirmed on CT scan (Figure 1). Esophagogastroduodenoscopy identified distal esophagitis presented with linear ulcerations covered with fibrin; gastric mucosa without changes; both in the esophagus and in the stomach hemorrhagic content; functional nonmechanical gastric outlet obstruction; distended duodenum with numerous shallow ulcerations covered with fibrin, without active bleeding (ischemic changes suspected), abundant blood reflux from distal parts of the
duodenum. Multislice CT angiography of abdomen excluded occlusion of the mesenteric vessels.

Conservative management with nasogastric suction, the usage of laxatives and prostigmin injections as well as colonic irrigation was performed with a good clinical response. In addition, fluid and electrolytes resustitation including blood products and gastroprotective medications was introduced. During the next several days melena was present.

During hospitalization, the control chest X-ray showed a pneumonic infiltration typical for aspiration bronchopneumonia. Despite the fact that antibiotic treatment had been introduced, the primarily detected process in the lungs was propagated. Because of the acute respiratory failure, the patient required mechanical ventilation for 10 days.

From the day 25, the patient was completely recovered and was discharged from the hospital at day 30.

DISCUSSION

The clinical picture of poisoning in our patient pointed to the dominant effects of clozapine overdose. To the best of our knowledge, gastrointestinal complications of acute clozapine poisoning in the drug-naïve patient with the onset in the phase of recovery, after reversal of typical nervous systems disturbances, have not been previously discussed.

In this report, clinical signs of clozapine toxicity (miotic pupils, hypersalivation, high puls rate) were predominated on admission. They were accompanied with high both CRP and serum CK levels. From the second day, benign fever was detected, while from the
seventh day gastrointestinal complications, including esophagitis, paralytic ileus and gastrointestinal bleeding, were noticed. In comparison to other second-generation antipsychotics, the occurrence of gastrointestinal hypomotility as side effect has been most widely reported for clozapine, presumably due to the anticholinergic activity [3, 5]. Moreover, taking into account the antidopaminergic effect of antipsychotics, the underlying mechanism of digestive hypoperfusion and ischemia could be suggested. Due to its DA1 vascular receptors agonistic activity and subsequent vasodilatatory effects at low dose, dopamine could improve the mesenteric perfusion. On the other hand, antipsychotics, as antidopaminergic agents, could play an additional role in digestive ischemia through inhibition of dopamine-dependant mesenteric vasodilatation [17]. The bowel ischemia related to motility disturbances could be also explained by untreated intestinal obstruction or pseudoobstruction, which can provoke not just gastrointestinal bleeding and distension, but also more serious complications such as gastrointestinal necrosis and perforation [18]. The authors agree that fever, abdominal tenderness, rigidity and leukocytosis should raise the possibility of ischemia, toxic megacolon, or perforation [6]. Considering clozapine-induced esophageal complications, Laker et al., in a series of 36 patients treated with clozapine, reported 4 cases who developed upper gastrointestinal symptoms suggestive of reflux oesophagitis within 6 weeks of starting this drug [19].

It is accepted that the acute toxicity of clozapine is much greater in the absence of tolerance to the drug [5, 20]. Hence, Shu-Chin Yu et al., by reporting rapid development of fatal bowel infarction within 1 week after clozapine treatment, raises the issue that the absence of tolerance to clozapine could cause mortality associated with clozapine-induced ischemic bowel at the beginning of the administration [21]. In the review of the French pharmacovigilance database, the time between the beginning of antipsychotics involved and onset of the digestive complications was variable – between 6 days and 20 years, with a median duration of 35 days [22].

Other medication ingested by our patients may also contribute to the emergence of ileus. Even though constipation is a minor side effect of haloperidol therapy, because of its weak anticholinergic effects, significant gastrointestinal hypomotility and rarely paralytic ileus can occur with biperiden [6, 23]. Clinicians should bear in mind that the anticholinergic interaction from the combination of these two drugs with clozapine may be responsible for serious impairment of gastrointestinal hypomotility. In the review by Peyrière et al, 65.8% of patients with antipsychotics-induced digestive complications were treated by at least one concomitant antimuscarinic medication beside clozapine [22].

Clozapine-induced benign fever is also a well-recognized adverse effect, and it is highly advisable for health care professionals to be cognizant of both agranulocytosis and neuroleptic malignant syndrome considering its differential diagnosis [24-26]. Thus, atypical neuroleptic malignant syndrome associated with atypical antipsychotics has been defined as rare small increases in CK, minor muscle rigidity and mild fever [27, 28]. In particular, Levenson’s definition of neuroleptic malignant syndrome appears the most flexible because it allows for its diagnosis to be made in the absence of either fever or rigidity [29, 30]. An inflammatory response with an acute phase reaction elevation in CRP and the subsequent fever are caused by the effect of clozapine on the cytokine system via IL-6 and TNF-α as it was postulated by the investigators [31]. Štuhec et al. and Kohen et al., in recently published case reports, discussed development of a benign fever in association with a rise in the CRP level in the patients on clozapine treatment. The fever resolved and CRP level declined progressively with clozapine discontinuation [32, 33].

CONCLUSION

Based on previously presented Clozapine Induced Delayed Ogilvie Syndrome at young 31 year old man, clinicians should be aware of the potential of atypical antipsychotic drugs to cause ileus, particularly in combination with other drugs with antimuscarinic properties, and ready to rapidly detect and treat intestinal atony thus preventing life-threatening complications. In addition, it should be highlighted that serum clozapine levels may not equate to clinical toxicity and the drug-naïve patient require more careful observation for complications in clozapine toxicity settings.
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CONFLICTS OF INTEREST

All authors declare no conflict of interest.

REFERENCES


Da li je klozapin odgovoran za odloženi Ogilvie sindrom i gastrointestinalno krvarenje u uslovima predoziranja?


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KRATAK SADRŽAJ

Uvod: Opisan je Ogilvie sindrom i gastrointestinalno krvarenje kao komplikacije nakon prestanka tipične kliničke slike akutnog trovanja klozapinom.


Zaključak: Kliničari treba da budu svesni da atipični antipsihotici imaju potencijal da prouzrokuju ileus, posebno u kombinaciji sa drugim antimuskarinskim lekoviima, i spemni da brzo prepoznaju i leče atoniju creva tako prevenirajući životno-ugrožavajuće komplikacije. Koncentracija klozapina u serumu ne mora korelirati sa kliničkom slikom i pacijenti koji nisu imali ranija iskustva sa lekom zahtevaju pažljiviju observaciju u sličaju trovanja klozapinom.

Ključne reči: atipični antipsihotici, klozapin, akutno trovanje, odložene komplikacije, Ogilvie sindrom, gastrointestinalno krvarenje