



CLOZAPINE- SAFE PRESCRIBING - WE ARE COUNTING ON YOU

- REGULARLY CHECK FOR SYMPTOMS OF NEUTROPENIA AND AGRANULOCYTOSIS
- MANAGE CONSTIPATION PROACTIVELY
- ASSESS FOR MYOCARDITIS AND CARDIOMYOPATHY
- BE AWARE OF OTHER ADVERSE REACTIONS AND INTERACTIONS ESPECIALLY SMOKING

Clozapine has drastically improved the lives of patients with treatment resistant schizophrenia, but it can cause serious adverse effects. Clozapine can only be initiated by a Psychiatrist, but symptoms of adverse effects may present first in Primary Care.

Clozapine is associated with neutropenia which may progress to a potentially fatal agranulocytosis. Severe constipation, myocarditis and adverse metabolic effects need to be assessed regularly; these are not dosedependent.

REGULARLY CHECK FOR SYMPTOMS OF NEUTROPENIA AND AGRANULOCYTOSIS

Advise patients to be alert for symptoms of neutropenia (eg fever, sore throat or flu-like symptoms). Anyone taking clozapine and presenting with these symptoms needs an urgent full blood count, medical review, and their mental health team notified immediately.

If you have a patient who is taking clozapine, please consider putting an alert on their file to raise awareness that even minor symptoms could be serious.

Regular blood monitoring is necessary for pharmacists to supply clozapine. Patients are registered with the manufacturer's blood monitoring database; *ClopineConnectTM* for Clopine® brand, or *CareLink PlusTM* for Clozaril® brand. General Practitioners who are involved in prescribing will need to use the blood monitoring database; they will be provided with training and access via a web portal.

A baseline full blood count is required 10 days prior to commencing treatment, then **weekly** full blood counts are needed during the first 18 weeks of treatment. 28 day ('monthly') monitoring is required thereafter and for 4 weeks after discontinuation. Despite this vigilance, deaths from agranulocytosis have occurred in New Zealand.⁴ Agranulocytosis tends to develop during the first 6 months of treatment and is not dose-related. Neutropenia can occur at any time.

Other medicines that can also increase the risk of neutropenia should not be used concurrently with clozapine. These include some antibiotics co-trimoxazole, trimethoprim, nitrofurantoin), carbamazepine, and antineoplastics that are associated with bone marrow suppression.

MANAGE CONSTIPATION PROACTIVELY

Clozapine causes constipation in at least 80% of patients and can occur at any time. Complications from constipation are the most common reason for clozapineinduced mortality and serious morbidity in New Zealand.

The risk is increased with concurrent anticholinergics (eg tricyclic antidepressants), opioids, iron supplements and calcium channel blockers, other illnesses and high doses.

Provide access to stimulant laxatives, give dietary advice and address pre-existing constipation prior to initiating clozapine. Bulking laxatives may make existing constipation worse.

Symptoms of serious complications include abdominal pain or distension, vomiting, watery diarrhoea (overflow), reduced appetite and nausea. Advise patients and their families to report these symptoms immediately; treatment interruption and review is recommended. Patients with no bowel movement in five days should be admitted to hospital for treatment.

ASSESS FOR MYOCARDITIS AND CARDIOMYOPATHY

Clozapine is associated with a small but significant risk of myocarditis and cardiomyopathy; fatalities have been reported in New Zealand. Symptoms of cardiomyopathy include signs of heart failure, flu-like illness, sinus tachycardia, hypotension and chest discomfort.

Symptoms of cardiomyopathy include signs of heart failure, flu-like illness, cough, fever, sinus tachycardia, hypotension and chest discomfort.

Symptoms of myocarditis are often non-specific and may include flu-like illnesses, gastrointestinal upset, unexplained fatigue, chest pain, dyspnoea, marked fluctuations in blood pressure, and electrocardiogram changes. Patients presenting with any of these symptoms

🛏 continued





CLOZAPINE

should be referred urgently for a cardiology review, and the patient's psychiatrist notified.

BE AWARE OF OTHER ADVERSE REACTIONS AND INTERACTIONS

Adverse effects including sedation and postural hypotension are more pronounced if the dose is not gradually titrated or a 'normal' dose is given after a treatment interruption. Weight gain and glucose intolerance (leading to type 2 diabetes) can also occur. Metabolic changes may increase cardiovascular and cerebrovascular risk. Monitor weight, blood pressure, HbA_{1c} and lipid parameters closely and encourage dietary and lifestyle modifications such as exercise. If necessary, use risk-lowering medications such as statins and metformin.

Other problematic side effects include enuresis, hypersalivation, increased sweating and tachycardia. Clozapine has been associated with increased incidence of pneumonia, and pulmonary aspiration, and can lower the seizure threshold.

Interactions

Clozapine interacts with a range of medicines that may decrease the plasma levels of clozapine. These include carbamazepine, phenytoin, rifampicin and omeprazole. Clozapine levels are affected by **cigarette smoking**, however it is the constituents of smoke, not nicotine itself, that increases the metabolism of clozapine. Clozapine levels can double when patients stop smoking. If patients stop smoking, monitor plasma clozapine levels and contact their mental health service; a dose reduction may be required. It is important that patients are aware of this and report any changes in smoking status during treatment.

Other medicines including erythromycin, ciprofloxacin and selective serotonin re-uptake inhibitors (SSRIs) such as paroxetine and fluoxetine can increase the plasma levels of clozapine. A high caffeine intake (more than 400mg/day) can increase levels, and subsequently decrease them by nearly 50% after a 5 day caffeinefree period.

Note: Clozapine can also enhance the central effects of alcohol, CNS depressants, and benzodiazepines.

REFERENCES

- McKean A. Safer prescribing of high-risk medicines: Clozapine. Best Practice Journal 2014;62:46-9 www.bpac.org.nz/BPJ/2014/July/docs/BPJ62-saferprescribing.pdf (Accessed 17-01-19)
- Clozapine: A reminder about safe and effective use. Best Practice Journal 2008;14:12-13. <u>www.bpac.org.nz/</u> <u>magazine/2008/june/docs/bpj14_clozapine_pages_12-</u> <u>13.pdf</u> (Accessed 17-01-19)
- 3. The New Zealand Formulary Clozapine <u>http://nzf.org.nz/</u> <u>nzf_2168</u> (Accessed 17-01-19)

ACKNOWLEDGEMENTS

We wish to thank Keith Crump and Ariel Hubbert, Mental Health Pharmacists at Waitematā District Health Board, for their valuable contribution to this bulletin.

CLICK HERE FOR FURTHER INFORMATION ON CLOZAPINE AND A FULL REFERENCE LIST

For further information on other high-risk medicines visit our website at : www.saferx.co.nz

No: 0182-01-035, Issued January 2019; Review: January 2022

DISCLAIMER: This information is provided to assist primary care health professionals with the use of prescribed medicines. Users of this information must always consider current best practice and use their clinical judgement with each patient. This information is not a substitute for individual clinical decision making. Issued by the Quality Use of Medicines Team at Waitematā District Health Board, email: feedback@saferx.co.nz