How to Provide Psychiatric Consultation for Patients on Clozapine

Learning Objectives:

- 1) Appreciate the role of CL psychiatrists in maintaining clozapine
- 2) Recognize pharmacokinetic factors that affect clozapine effectiveness and risk of toxicity
- 3) Learn strategies to prevent and manage adverse effects of clozapine

Step 1: Appreciate the unique role of clozapine

- Clozapine is effective for treatment-refractory schizophrenia, aggression in dementia, and psychosis in Parkinson and related disorders.
 - Ongoing need for clozapine may be questioned in cases of sedation, delirium, pneumonia, seizures, and other adverse effects.
 - Consulting for patients on clozapine involves educating about clozapine being more
 effective than other antipsychotics in specific cases and about strategies for
 managing treatment-limiting adverse effects.
 - Determine the patient's last dose of clozapine as soon as possible. If clozapine has been missed for more than 48 hours, clozapine will need to be gradually re-titrated. Education about swift re-initiation of clozapine at admission to avoid re-titration can prevent clinical relapse. See below for recommended re-initiation if doses have been missed.

Step 2: Learn basic clinical pharmacology of clozapine

- 95% protein bound; half-life 4-8 hours; extensive first pass effect; 45-65% bioavailable; available as a tablet and as an oral suspension
- Clozapine is typically initiated at 25 mg PO QHS nightly and maintained at this dose for 3-4 days. It is conservatively titrated as follows: 50 mg nightly for 3-4 days □ further increases of 25 mg twice weekly until a target dose of 300 mg is reached (100 mg in elderly). Levels should then be checked and dose adjusted based on clinical response and drug levels (see below)
 - Standard daily dose of clozapine: 300-600 mg/day; maximum daily dose of 900 mg/day.
 - i. In elderly patients target dosing is 100-200 mg/day
 - ii. In Parkinson and related disorders target dose is approximately 50 mg/day
 - o Dangers of rapid titration include sedation, hypotension, seizures, agranulocytosis, neuroleptic malignant syndrome, mucus plugs and subsequent respiratory collapse
 - o Initial dose and speed of titration can be adjusted based on risk factors for adverse effects, tolerability and ability to monitor the patient. For example, an initiation dose of 6.25 mg or 12.5 mg nightly may be trialed in elderly patients with hypotension rather than 25 mg nightly.
- Therapeutic drug monitoring optimizes effectiveness and prevents adverse events. A trough steady-state concentration of 350–600 µg/L is recommended in schizophrenia.
- Re-initiation after lapse in treatment: board recommendation for re-starting Clozapine after various gaps in length are as follows: (1)



- O Up to 48 hours: no re-titration required
- 48-72 hours: Restart with half of the previous prescribed total daily dose on day one
 (in divided does 12 hours apart). Next give 75% of previous daily dose on day two
 and, if tolerated, the whole of the previous daily dose in the normal dosing schedule
 on day three.
- 72 hours one week: Begins with 12.5mg once or twice daily. Increase according to tolerability over 3 days
- o More than one week: titrate as if a new patient.
- Clozapine is primarily metabolized in the liver by the cytochrome P450 (CYP P450) isoenzyme 1A2 into norclozapine
 - Systemic inflammation, infection [e.g., COVID-19 (2)], and nicotine discontinuation can decrease CYP1A2 activity potentially leading to higher levels and possible clozapine toxicity
 - In COVID-19 and other respiratory infections with a systemic inflammatory response (evidenced by fever, increase in CRP) decrease dose of clozapine by at least 50%
 - ii. Tobacco induces CYP1A2 and increases metabolism of clozapine; abstaining from smoking in the hospital \(\precedeta \) decreases in clozapine metabolism \(\precedeta \) potentially leading to higher levels/toxicity; consider decreasing dose of clozapine during hospitalization.
 - Some East Asians may be slow metabolizers of clozapine (data from Korean,
 Chinese, Vietnamese populations). Lower doses than listed above are effective.
 - o Important drug interactions with clozapine are listed in Table 1.

Step 3: Describe adverse effects of clozapine and their management

- Clozapine has many serious and non-serious adverse effects. The FDA provides "black box" warnings for severe neutropenia, orthostatic hypotension, bradycardia, seizure, syncope, myocarditis, cardiomyopathy, and mitral valve incompetence, as well as increased mortality in older adults with dementia-related psychosis.
- The greatest risk of death comes from agranulocytosis, myocarditis/cardiomyopathy and pulmonary embolism/DVT. (3)
- If a decision is made to discontinue clozapine, it should not be abruptly discontinued (unless myocarditis or severe neutropenia are present); it should be gradually tapered over 1-2 weeks. Sudden discontinuation can lead to exacerbation of underlying psychiatric illness, clozapine withdrawal catatonia, and cholinergic rebound
 - Treatment of clozapine withdrawal catatonia includes resuming clozapine in addition to standard treatment for catatonia. Even low doses of clozapine can reverse the course of a clozapine-withdrawal catatonia.
- The prevention and management of common adverse effects encountered by CL psychiatrists are described in Table 2.



References

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Table 1: Important drug interactions with clozapine (2)

Medications/Substances	Effects
Cigarette smoking Carbamazepine, rifampin, phenytoin, phenobarbital, St. John's Wort	May ↓ reduce clozapine levels via CYP 1A2 or 3A4 induction resulting in decreased efficacy of clozapine
Fluvoxamine, ciprofloxacin, oral contraceptives, caffeine antidepressants: (fluoxetine, escitalopram, paroxetine, duloxetine, sertraline), antibiotics: (erythromycin), cimetidine, quinidine, terbenafine	May ↑ increase clozapine levels via CYP 1A2 or 2D6 inhibition leading to adverse reactions
Anticholinergics (e.g., benztropine, diphenhydramine, hydroxyzine)	Additive anticholinergic activity. Monitor closely for significant anticholinergic adverse effects (e.g., constipation, hypotension).
Lithium	Combined use is associated with neurotoxicity (delirium, movement disorders, myoclonus, and seizures)
Bone marrow suppressants (e.g., antineoplastics)	Caution due to increased risk and/or severity of bone marrow suppression.
CNS depressants and general anesthesia (e.g., alcohol, opioids, benzos)	Caution because of additive CNS depressant effects including respiratory effects
Medications that lower seizure threshold (e.g., bupropion)	Extreme caution when coadministering with clozapine due to increased risk of seizures
Highly protein bound drugs (e.g., warfarin, digoxin)	Clozapine may increase levels of protein bound drugs and vice versa. Adjust dose if necessary.



Table 2: Prevention, and management of clozapine related adverse effects

	Description	Prevention/Monitoring	Management
Sialorrhea	Increases risk of aspiration pneumonia	None	Cover pillow with towel
	Stigmatizing		Elevate head of bed and encourage
	Thought to be due to muscarinic-4 antagonism,		sleeping on side reduces risk of
	alpha-2 antagonism with unopposed beta		aspiration
	agonism, decreased laryngeal peristalsis,		Ipratropium oral spray 0.06% up to
	decreased swallowing via reduction of GI		3 sprays TID □ Ophthalmic atropine
	motility		1% up to 2 drops TID sublingually
			☐ Oral glycopyrrolate 2-4 mg PO
			QHS □ Terazosin 1-2
			mg PO QHS
			Botox in refractory cases
			·

	Description	Prevention/Monitoring	Management
Constipation, ileus, bowel perforation	50-80% prevalence of constipation Due to anticholinergic and antiadrenergic effects Risk factors: high dose, high level, concomitant administration with other anticholinergic medications Presentation: constipation, abdominal distension, nausea, emesis Diagnosis: subjective report, exam, abdominal X-ray, CT of abdomen	Patients with schizophrenia may not report symptoms due to higher pain thresholds and negative symptoms. Regular inquiry into bowel function is necessary. Hydration High fiber diet Exercise Start stool softeners or laxatives in patients at high risk Docusate polyethylene glycol sennakot or dulcolax lubiprostone	Constipation and ileus: Docusate polyethylene glycol sennakot or dulcolax magnesium citrate/other enemas/manual disimpaction Obstruction: nasogastric suction; surgery may sometimes be necessary
Clozapine toxicity	Delirium, neutropenia, orthostatic hypotension, tachycardia, ileus, urinary retention, mydriasis/miosis, hypersalivation, and seizures.	Recognize of drug-drug and drug disease interactions that can lead to elevated clozapine levels and clozapine toxicity; prophylactically decrease clozapine doses	Decrease or hold dose of clozapine. Check level. May take several days for results to be available but level aids in confirming suspected clozapine toxicity)



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Seizures	Prevalence of 3.5-8 % Tonic-clonic seizures are most common Risk factors: Epilepsy High doses (doses > 600 mg/day) (3) Rapid titration during initiation phase	Prophylaxis with anticonvulsants for patients with: Seizure disorder myoclonic jerks (may be precursor to generalized tonic-clonic seizures for patients on clozapine) Paroxysmal spike/sharp wave discharges Clozapine plasma level > 500 ug/L Clozapine dose > 500 mg/day	Obtain EEG Neurology consultation Add anticonvulsant (e.g., valproic acid) Clozapine rarely needs to be discontinued		

	Description	Prevention/Monitoring		Management	
	Rare, idiosyncratic reaction	Severity		Monitoring	Management
Neutropenia	Makes body vulnerable to bacterial infections. Frequently present with fever, headache, cough, sore throat, bleeding gums or sepsis The Clozapine REMS program provides centralized clozapine access to minimize the risk of clozapine-associated severe neutropenia	Normal	ANC >1500/mcL	Weekly from initiation to 6 months Every 2 weeks from 6 to 12 months Monthly after 12 months	Continue treatment
	Benign ethnic neutropenia (BEN) is common in people of African or Eastern Mediterranean heritage. Alternate monitoring parameters exist for those with established	Mild	ANC 10001499/mc L	3x/week monitoring until ANC ≥ 1500/µl	Continue treatment
	BEN For patients who have been on long-term clozapine without neutropenia and have new onset neutropenia with other constitutional symptoms, premature attribution of neutropenia to clozapine should be avoided.	Moderate	ANC 500999/mcL	Daily until ANC ≥ 1000/μ1 then, 3x/week until ANC ≥ 1500/μ1	Interrupt treatment for suspected clozapine induced neutropenia Hematology consultation Consider filgrastim or lithium



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To this end, a broad differential diagnosis and thorough work-up for neutropenia is warranted.	Severe	ANC < 500/mcL	Daily until ANC ≥ 1000/µl then, 3x/week	Interrupt treatment for suspected clozapine induced neutropenia Transfer to hospital Place in reverse isolation Administer filgrastim Aggressively work up signs and symptoms Report to Clozapine REMS
	should ideally be order to do so Cl Multidisciplinary parameters, thres blood draws Consider initiation	terfere with cance continued if the pozapine REMs pro team discussion shold for disconting on of filgrastim du	patient is well establi ogram will need to be with heme-onc regar muation of clozapine,	ding alternate monitoring and increased frequency of lir of chemotherapy to support

	Description	Prevention/Monitoring	Management
Myocarditis	1-3% prevalence High mortality (10 – 30%) Typically seen in first 4 weeks post-initiation Most likely IgE mediated hypersensitivity reaction Presentation: chest pain, palpitations, dyspnea, tachycardia Diagnosis: History and exam along with CRP, cardiac enzymes, eosinophil count, ECG findings, CXR, cardiac echo (5)	Monitor vitals, cardiac enzymes, and CRP weekly during first 4 weeks of titration Obtain baseline ECG	Discontinue clozapine and admit to intensive care unit Supportive use of medications to improve cardiac functioning and minimize the risk of cardiac failure such as diuretics, beta-blockers, and ACE inhibitors, corticosteroids

