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SA Health

Cancer Chemotherapy Rapid Desensitisation Schedule: OXALIPLATIN

> SA Health Cancer Drug Committee July 2023



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Applicant Details

Consultant Name: Sid Selva Position: medical Oncologist		
Clinical Unit, Hospital/LHN: Cancer Services / RAH /CALHN		
Telephone:	Pager:	
Mobile:	Email:	

Supporting Tumour Stream Lead Details

Consultant Name: SAHCDC INIT Position:	IATED	
Clinical Unit, Hospital/LHN:		
Telephone:	Pager:	
Mobile:	Email:	

Supporting Specialist Pharmacist Details

Name: Helen Martin Position: Senior Pharmacist, Oncology	
Clinical Unit, Hospital/LHN: SALHN/FMC	
Telephone: 82046286	Pager: 38854
Mobile:	Email: helen.martin@sa.gov.au

Supporting Specialist Nurse Details

Name: Position:	
Clinical Unit, Hospital/LHN:	
Telephone:	Pager:
Mobile:	Email:

SA Health Cancer Drug Committee Use only:

Application received (date):				
Confirmation of costing confi	rmed* □			
Approval Status				
Conditions of approval (if any	/):			
REJECTED 🗆				
Reason(s) for rejection:	Reason(s) for rejection:			
Treatment Risk Level allocat	ed:			
SAH-CDC comments (if any				
and confirm the decision ma	n and to the best of my knowledge the information contained within is correct de by the SA Health Cancer drug Committee in submitting this protocol to the SA emotherapy Protocol Register:			
SAH-CDC Chair (or delegate	e): Position:			
Signature:	Date:			
Protocol Name				
Protocol Number				

Rapid Desensitisation Schedule-Oxaliplatin

Treatment Schedule - Summary

This treatment schedule describes a method of drug administration which allows the safe re-introduction of oxaliplatin after a previous hypersensitivity reaction. Oxaliplatin is administered in an incremental fashion (with increased premedication and at gradually increasing concentrations and infusion rates), until the full therapeutic dose has been given. The state of tolerance induced is temporary and disappears once oxaliplatin is cleared. The patient remains hypersensitive to oxaliplatin. Therefore, this schedule must be followed for every dose of oxaliplatin in the approved cancer chemotherapy protocol, until the patient has completed all treatment cycles.

This treatment schedule for desensitisation must be used in conjunction with the approved oxaliplatin-containing cancer chemotherapy protocol.

Drug	Dose	Route	Day
Oxaliplatin	As per approved protocol	IV	As per approved protocol

Risk Rating High

First desensitisation must be administered as an inpatient. If there is no hypersensitivity reaction during this session, subsequent cycles may be administered in the outpatient setting.

Indications and Patient Population

Patients who have experienced moderate-to-severe hypersensitivity reactions (including severe anaphylaxis) to oxaliplatin, for whom alternative treatment of equal efficacy or toxicity is not available.

Contraindications and Precautions

Patients who have experienced severe life-threatening immunocytotoxic reactions, vasculitis or exfoliative skin diseases such as Stevens-Johnson Syndrome/Toxic Epidermal Necrolysis (SJS/TEN), or Drug Reactions with Eosinophilia and Systemic Symptoms (DRESS).

This treatment schedule is not recommended for unstable patients (eg uncontrolled asthma, cardiac disease, or haemodynamic instability); however, desensitisation may be considered once these underlying conditions have stabilised.

Treatment S	Freatment Schedule - Detailed				
Drug	Dose		Step	Rate of Administration	
Oxaliplatin	Bag 1:			1.	2 mL/h for 15 minutes, then
-	1/100 dilution in 250 mL glucose		Э	2.	5 mL/h for 15 minutes, then
	5% (discard after 1 hour)			3.	10 mL/h for 15 minutes, then
				4.	20 mL/h for 15 minutes.
	Bag 2:			5.	5 mL/h for 15 minutes, then
	1/10 dilution in 250 mL glucose 5		5%	6.	10 mL/h for 15 minutes, then
	(discar	d after 1 hour)		7.	20 mL/h for 15 minutes, then
				8.	40 mL/h for 15 minutes.
	Bag 3:			9.	10 mL/h for 15 minutes, then
	100% (concentration in 250 mL		10.	20 mL/h for 15 minutes, then
	glucose	e 5%		11.	40 mL/h for 15 minutes, then
	(Pharn	nacist to calculate final		12.	75 mL/h to complete prescribed
	volum	e to be infused to delive	ər		dose, then discard any
	100% o	00% of dose)			remaining solution.
Desensitisatio	esensitisation premedication * Drug Dose Frequency / timing				/ / timing
Dexametha	sone	8 mg PO	13	13 hours prior to oxaliplatin	
Cetirizine		10 mg PO	13	hours p	rior to oxaliplatin
Famotidine		20 mg PO	13	hours p	rior to oxaliplatin
Dexametha	sone	8 mg PO	7 h	7 hours prior to oxaliplatin	
Cetirizine		10 mg PO	7 h	ours pri	or to oxaliplatin
Famotidine		20 mg PO	7 h	7 hours prior to oxaliplatin	
Dexametha	sone	20 mg IV	60	60 minutes prior to oxaliplatin	
Cetirizine		10 mg PO		60 minutes prior to oxaliplatin	
Famotidine		20 mg PO	60 minutes prior to oxaliplatin		
For breakthrough hypersensitivity reactions:					
Hydrocortis	sone	100 mg IV PRN	At commencement of bag 3 (100 % concentration) of oxaliplatin		
Cetirizine		10 mg PO PRN		At commencement of bag 3 (100 % concentration) of oxaliplatin	
* see approved	d cance	er chemotherapy protoco	for	oremedi	cation required for other drugs

administered as part of the protocol (to avoid duplication)

Monitoring (additional to approved protocol)

Observe the patient during administration and for 2 hours after the completion of bag 3 of oxaliplatin for any sign of breakthrough hypersensitivity reactions.

If a hypersensitivity reaction occurs, stop the infusion and treat according to institutional protocols. After the reaction has subsided, the infusion can be restarted from the step at which it had been paused. If repeated or severe reactions occur, the treatment schedule can be further adjusted by prolonging the step before the reaction occurred, adding an additional (dilution) step, and/or administering prophylactic medication before the step at which the patient had a reaction. This must be done in consultation with the treating consultant and oncology/haematology pharmacist.

Supporting Documents

Seghers S, Teuwen LA, Beyens M, et al. Immediate hypersensitivity reactions to antineoplastic agents - A practical guide for the oncologist. *Cancer Treat Rev.* 2023;116:102559. <u>https://doi:10.1016/j.ctrv.2023.102559</u>

Pagani M, Bavbek S, Alvarez-Cuesta E, et al. Hypersenstivity reactions to chemotherapy: an EAACI Position Paper. *Allergy*. 2022;77:388–403. <u>https://doi.org/10.1111/all.15113</u>

Tsao LR, Young FD, Otani IM, Castells MC. Hypersensitivity Reactions to Platinum Agents and Taxanes. *Clin Rev Allergy Immunol*. 2022;62(3):432-448. <u>https://doi:10.1007/s12016-021-08877-y</u>

Caiado J, Castells MC. Drug Desensitizations for Chemotherapy: Safety and Efficacy in Preventing Anaphylaxis. *Curr Allergy Asthma Rep.* 2021;21(6):37. Published 2021 Jul 7. https://doi:10.1007/s11882-021-01014-x

Lee CW, Matulonis UA, Castells MC. Rapid inpatient/outpatient desensitization for chemotherapy hypersensitivity: standard protocol effective in 57 patients for 255 courses. *Gynecol Oncol.* 2005;99(2):393-399. <u>https://doi:10.1016/j.ygyno.2005.06.028</u>

Parel M, Ranchon F, Nosbaum A, et al. Hypersensitivity to oxaliplatin: clinical features and risk factors. *BMC Pharmacol Toxicol*. 2014;15:1. Published 2014 Jan 13. <u>https://doi:10.1186/2050-6511-15-1</u>

Castells M. Rapid desensitization of hypersensitivity reactions to chemotherapy agents. *Curr Drug Saf.* 2006;1(3):243-251. <u>https://doi:10.2174/157488606777934413</u>

Hypersensitivity reaction. eviQ clinical resources. https://www.eviq.org.au/



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For more information

Clinical System Support and Improvement Division Office of the Chief Pharmacist Department for Health and Wellbeing, SA Health Level 8, Citi Centre 11 Hindmarsh Square Adelaide, SA 5000 Telephone: +61 8 8204 1944 www.sahealth.sa.gov.au



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