RS2125 - Tocilizumab

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	Name:
	NHI:
	14111.
treatment of acute lymphoblastic leukaemia Tocilizumab is to be administered at doses no greater to f 12 mg/kg) The patient is enrolled in the Malaghan Institute of Med The patient has developed CRS or Immune Effector Cetherapy for the treatment of relapsed or refractory B-ce Tocilizumab is to be administered according to the constitution.	ell-Associated Neurotoxicity Syndrome (ICANS) following CAR T-Cell
evious use equired after 6 months ck boxes where appropriate) ped by, or recommended by any relevant practitioner, or in a pital. atient was being treated with tocilizumab prior to 1 Februar Rheumatoid arthritis Systemic juvenile idiopathic arthritis Adult-onset Still's disease Polyarticular juvenile idiopathic arthritis	occordance with a protocol or guideline that has been endorsed by the Health y 2019
	The patient has developed grade 3 or 4 cytokine release treatment of acute lymphoblastic leukaemia Tocilizumab is to be administered at doses no greater to of 12 mg/kg) The patient is enrolled in the Malaghan Institute of Med The patient has developed CRS or Immune Effector Cetherapy for the treatment of relapsed or refractory B-cetherapy for the treatment of relapsed or refractory B-cetherapy for the treatment of a maximum of 3 doses Vious use quired after 6 months k boxes where appropriate) ed by, or recommended by any relevant practitioner, or in a sital. The patient has developed CRS or Immune Effector Cetherapy for the treatment of relapsed or refractory B-cetherapy for the treatment of the maximum of 3 doses

I confirm that the above details are correct:		
Signed:	Date:	

June 2025

PRESCRIBER				P	PATIENT:											
Name:														Ν	lame:	
Ward:														Ν	IHI:	
Tocilizu	ıma	ab	- con	tinue	ed											
INITIATI Re-asse								nts p	reviou	ısly tr	reated	d with a	adali	lim	umab or etanercept)	
Prerequ								te)								
O and												Practitio Ilth NZ I			the recommendation of a rheumatologist, or in accordance with a al.	
an)	The p	atie	nt has	had	an initia	al Spe	cial Au	uthorit	ty app	oroval fo	or ad	ilat	imumab and/or etanercept for rheumatoid arthritis	
	The patient has experienced intolerable side effects from adalimumab and/or etanercept The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or etanercept such that they do not meet the renewal criteria for rheumatoid arthritis															
an	and						_ 									
		or		The	patie	nt is	serone	gative	for bot	th anti	ti-cycli	ic citrul	llinate	ed	peptide (CCP) antibodies and rheumatoid factor	
			an	O	The	pat	ient has	been	starte	ed on r	rituxin	nab for	rheu	um	atoid arthritis in a Health NZ Hospital	
				O	C	A	four m	onths	followi	ing the	e initia	al cours	se of	f rit	uximab the patient has received insufficient benefit such that they darthritis	
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Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRES	CRIB	ER	PATIENT:	PATIENT:					
Name	:		Name:	Name:					
Ward:	rd:NHI:								
Tocil	izum	nab	- continued						
Re-a	ssessi	ment	Rheumatoid Arthritis at required after 6 months (tick boxes where appropriate)						
and			cribed by, or recommended by a rheumatologist or Practitioner on the recommendation of a rhe col or guideline that has been endorsed by the Health NZ Hospital.	umatologist, or in accordance with a					
	(and		Patient has had severe and active erosive rheumatoid arthritis (either confirmed by radiology in citrullinated peptide (CCP) antibody positive) for six months duration or longer	maging, or the patient is cyclic					
	and	C	Tocilizumab is to be used as monotherapy						
		or	Treatment with methotrexate is contraindicated						
	and		Patient has tried and did not tolerate oral and/or parenteral methotrexate						
	and	or	O Patient has tried and not responded to at least three months therapy at the maximum tol combination with another agent	lerated dose of ciclosporin alone or in					
		O.	O Patient has tried and not responded to at least three months therapy at the maximum tol combination with another agent	lerated dose of leflunomide alone or in					
	and								
		or	O Patient has persistent symptoms of poorly controlled and active disease in at least 20 ac						
			O Patient has persistent symptoms of poorly controlled and active disease in at least four a elbow, knee, ankle, and either shoulder or hip	active joints from the following: wrist,					
	and								
		or	Patient has a C-reactive protein level greater than 15 mg/L measured no more than one application	e month prior to the date of this					
		0.	O C-reactive protein levels not measured as patient is currently receiving prednisone thera day and has done so for more than three months	py at a dose of greater than 5 mg per					
Re-a	ssessi	ment	systemic juvenile idiopathic arthritis It required after 6 months (tick boxes where appropriate)						
(and			cribed by, or recommended by a rheumatologist or Practitioner on the recommendation of a rhe col or guideline that has been endorsed by the Health NZ Hospital.	umatologist, or in accordance with a					
	and	C	Patient diagnosed with systemic juvenile idiopathic arthritis						
	(C	Patient has tried and not responded to a reasonable trial of all of the following, either alone or methotrexate; non-steroidal anti-inflammatory drugs (NSAIDs); and systemic corticosteroids	in combination: oral or parenteral					

I confirm that the above details are correct:

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

PRESCR	IBER		PATIENT:				
Name:							
Ward:			NHI:				
Tocilizu	ımab	- COI	ntinued				
Re-asse	ssmer isites Pres	it requ (tick b	by, or recommended by a rheumatologist or Practitioner on the recommendation of a rheumatologist, or in accordance with a				
and	proto	col or	guideline that has been endorsed by the Health NZ Hospital.				
		 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for adult-onset Still's disease (AOSD) The patient has been started on tocilizumab for AOSD in a Health NZ Hospital 					
	an	or	O The patient has experienced intolerable side effects from adalimumab and/or etanercept O The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or etanercept such that they do not meet the renewal criteria for AOSD				
or							
	Patient diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 1992;19:424-430) Patient has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal antiinflammatory drugs (NSAIDs) and methotrexate Patient has persistent symptoms of disabling poorly controlled and active disease						
Re-asse Prerequ	ssmer isites Pres	it requ (tick b cribed	rticular juvenile idiopathic arthritis ired after 4 months boxes where appropriate) by, or recommended by a rheumatologist or Practitioner on the recommendation of a rheumatologist, or in accordance with a guideline that has been endorsed by the Health NZ Hospital.				
and	an	O	The patient has had an initial Special Authority approval for both etanercept and adalimumab for polyarticular course juvenile idiopathic arthritis (JIA) The patient has experienced intolerable side effects, or has received insufficient benefit from, both etanercept and adalimumab				
or	an an		Treatment with a tumour necrosis factor alpha inhibitor is contraindicated Patient has had polyarticular course JIA for 6 months duration or longer To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance O At least 5 active joints and at least 3 joints with limited range of motion, pain or tenderness after a 3-month trial of methotrexate (at the maximum tolerated dose) O Moderate or high disease activity (cJADAS10 score of at least 2.5) after a 3-month trial of methotrexate (at the				
		or	maximum tolerated dose) C Low disease activity (cJADAS10 score between 1.1 and 2.5) after a 6-month trial of methotrexate				

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PRESCRIBER	PATIENT:							
Name:	Name:							
Ward:	NHI:							
Tocilizumab - continued								
INITIATION – idiopathic multicentric Castleman's disease Re-assessment required after 6 months Prerequisites (tick boxes where appropriate)								
Prescribed by, or recommended by a haematologist, rheumatologist or in accordance with a protocol or guideline that has been endorse and	t or Practitioner on the recommendation of a haematologist or rheumatologist, ed by the Health NZ Hospital.							
Patient has severe HHV-8 negative idiopathic multicentric Ca	stleman's disease							
Treatment with an adequate trial of corticosteroids has prover and	n ineffective							
O Tocilizumab to be administered at doses no greater than 8 m	g/kg IV every 3-4 weeks							
INITIATION – moderate to severe COVID-19 Re-assessment required after 1 dose Prerequisites (tick boxes where appropriate)								
O Patient has confirmed (or probable) COVID-19								
Oxygen saturation of < 92% on room air, or requiring supplemental oxygen								
	O Patient is receiving adjunct systemic corticosteroids, or systemic corticosteroids are contraindicated							
Tocilizumab is to be administered at doses no greater than 8i	ng/kg IV for a maximum of one dose							
O Tocilizumab is not to be administered in combination with bar	citinib							
CONTINUATION – Rheumatoid Arthritis								
Re-assessment required after 6 months								
Prerequisites (tick boxes where appropriate)								
protocol or guideline that has been endorsed by the Health NZ Hos	on the recommendation of a rheumatologist, or in accordance with a pital.							
Following 6 months' initial treatment, the patient has at least significant response to treatment in the opinion of the physici or	a 50% decrease in active joint count from baseline and a clinically an							
On subsequent reapplications, the patient demonstrates at le a clinically significant response to treatment in the opinion of	ast a continuing 30% improvement in active joint count from baseline and the physician							
CONTINUATION – systemic juvenile idiopathic arthritis Re-assessment required after 6 months Prerequisites (tick boxes where appropriate)								
O Prescribed by, or recommended by a rheumatologist or Practitioner protocol or guideline that has been endorsed by the Health NZ Hos	on the recommendation of a rheumatologist, or in accordance with a pital.							
Following up to 6 months' initial treatment, the patient has ac improvement criteria (ACR Pedi 30) response from baseline	hieved at least an American College of Rheumatology paediatric 30%							
On subsequent reapplications, the patient demonstrates at le	ast a continuing ACR Pedi 30 response from baseline							

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ogist, or in accordance with a exate is limited by toxicity or t count and an improvement in active joint count and
of a haematologist or rheumatologist,
nat has been endorsed by the Health expoint inhibitor treatment for
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PRESCRIBER	PATIENT:				
Name:	Name:				
Ward:	NHI:				
Tocilizumab - continued					
CONTINUATION – immune checkpoint inhibitor toxicity in malignancy* Re-assessment required after 4 months Prerequisites (tick boxes where appropriate) Prescribed by, or recommended by any relevant practitioner, or in ac	ccordance with a protocol or guideline that has been endorsed by the Health				
The individual has shown clinical improvement and ongoing treatment is required Tocilizumab is to be administered at a maximum dose of 8 mg/kg fortnightly					
Note: Indications marked with * are unapproved indications.					