HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

July 2024

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.

PRESCRI	RIBER	PATIENT: Name:	
Name:			
Ward:		NHI:	
Secukin	numab		
Re-asses	ION – severe chronic plaque psoriasis, second-line biologic essment required after 4 months sisites (tick boxes where appropriate)		
and	Prescribed by, or recommended by a dermatologist, or in accorda Hospital.	nce with a protocol or guideline that has been endorsed by the Health NZ	
and	Hospital, for severe chronic plaque psoriasis	adalimumab or etanercept, or has trialled infliximab in a Health NZ	
	O The patient has experienced intolerable side effects for O The patient has received insufficient benefit from ada		
and	A Psoriasis Area and Severity Index (PASI) assessment or for at least the most recent prior treatment course, preferab each prior treatment course	Dermatology Quality of Life Index (DLQI) assessment has been completed by while still on treatment but no longer than 1 month following cessation of	
	The most recent PASI or DQLI assessment is no more than	1 month old at the time of application	
Re-asses	AUATION – severe chronic plaque psoriasis, second-line biologessment required after 6 months uisites (tick boxes where appropriate) Prescribed by, or recommended by a dermatologist, or in accordate Hospital.	nce with a protocol or guideline that has been endorsed by the Health NZ	
	O Patient's PASI score has reduced by 75% or more (Page 1)	ASI 75) as compared to baseline PASI prior to commencing secukinumab	
		I) improvement of 5 or more, as compared to baseline DLQI prior to	
and	nd _		

I confirm that the above details are correct:

Signed: Date:

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PRESCRIBER			PATIENT:			
Name:						
Ward				NHI:		
Secu	ıkinu	mak) - c	continued		
Re-a	ssessi equisi P	nent tes (t rescri	requ ick b ibed	e chronic plaque psoriasis, first-line biologic ired after 4 months poxes where appropriate) by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ		
and		ospita	al.			
		or	0	Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis		
				Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis		
	and			nt has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the ving (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin		
	and			SI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior ment course, preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course		
	() 1	Γhe r	most recent PASI or DQLI assessment is no more than 1 month old at the time of application		
psori recei and s	Note: A treatment course is defined as a minimum of 12 weeks of treatment. "Inadequate response" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the face, hand or foot, at least 2 of the 3 PASI symptom sub scores for erythema, thickness and scaling are rated as severe or very severe, and the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment.					
CONTINUATION – severe chronic plaque psoriasis, first-line biologic Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) Prescribed by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health Na Hospital.						
			O	Patient's PASI score has reduced by 75% or more (PASI 75) as compared to baseline PASI prior to commencing secukinumab		
		or	0	Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing secukinumab		
	and () s	Secu	kinumab to be administered at a maximum dose of 300 mg monthly		
Re-a	ssessı	nent	requ	osing spondylitis, second-line biologic ired after 3 months oxes where appropriate)		
and		rescri ospita		by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ		
	and) ₁	Гһе р	patient has had an initial Special Authority approval for adalimumab and/or etanercept for ankylosing spondylitis		
		or	O	The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept		
			O	Following 12 weeks of adalimumab and/or etanercept treatment, the patient did not meet the renewal criteria for adalimumab and/or etanercept for ankylosing spondylitis		

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PRESCRI	IBER	PATIENT:	
Name:		Name:	
Ward:		NHI:	
Secukin	numab	continued	
Re-asses	ssment i	ankylosing spondylitis, second-line biologic uired after 6 months boxes where appropriate)	
O and	Prescri Hospita	by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ	
and	_ b	owing 12 weeks initial treatment of secukinumab treatment, BASDAI has improved by 4 or more points from pre-secukinumab eline on a 10 point scale, or by 50%, whichever is less	
and	OF	sician considers that the patient has benefitted from treatment and that continued treatment is appropriate	
	<u></u>	ukinumab to be administered at doses no greater than 150 mg monthly	
Re-asses	ssment i	atic arthritis uired after 6 months coxes where appropriate) I by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ	
	and	Patient has had an initial Special Authority approval for adalimumab, etanercept or infliximab for psoriatic arthritis O Patient has experienced intolerable side effects from adalimumab, etanercept or infliximab Patient has received insufficient benefit from adalimumab, etanercept or infliximab to meet the renewal criteria for adalimumab, etanercept or infliximab for psoriatic arthritis	
or	and and	Patient has had severe active psoriatic arthritis for six months duration or longer Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose Patient has tried and not responded to at least three months of sulfasalazine at a dose of at least 2 g per day or leflunomide at a dose of up to 20 mg daily (or maximum tolerated doses) O Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen, tender joints O Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip O Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application O Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months	

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Name:					Name:		
Ward	:				NHI:		
Secu	ıkinı	ıma	b - c	ontinued			
Re-a	CONTINUATION – psoriatic arthritis Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) Orescribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.						
		or	0	clinically significant response to treatment in the opinion	ovement in active joint count from baseline and a clinically significant		
Secukinumab to be administered at doses no greater than 300 mg monthly				mg monthly			