## HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

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.

PRESCRIBER	PATIENT:					
Name:	Name:					
Ward:	NHI:					
Secukinumab Secukinumab						
INITIATION – severe chronic plaque psoriasis, second-line biologic Re-assessment required after 4 months						
Prerequisites (tick boxes where appropriate)						
Prescribed by, or recommended by a dermatologist, or in accordance Hospital.	ce with a protocol or guideline that has been endorsed by the Health NZ					
	dalimumab or etanercept, or has trialled infliximab in a Health NZ					
O The patient has experienced intolerable side effects fro	m adalimumab, etanercept or infliximab					
The patient has received insufficient benefit from adalin	numab, etanercept or infliximab					
for at least the most recent prior treatment course, preferably each prior treatment course	ermatology Quality of Life Index (DLQI) assessment has been completed 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					
CONTINUATION – severe chronic plaque psoriasis, second-line biologic Re-assessment required after 6 months						
Prerequisites (tick boxes where appropriate)						
Prescribed by, or recommended by a dermatologist, or in accordance Hospital.	be with a protocol or guideline that has been endorsed by the Health NZ					
	GI 75) as compared to baseline PASI prior to commencing secukinumab					
	improvement of 5 or more, as compared to baseline DLQI prior to					
Secukinumab to be administered at a maximum dose of 300	mg monthly					

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

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March 2025 RESTRICTIONS CRECKEST

Schedule. For community funding, see the Special Authority Criteria. **PRESCRIBER** PATIENT: Name: ..... Name: ..... Ward: ..... NHI: ..... Secukinumab - continued INITIATION - severe chronic plaque psoriasis, first-line biologic Re-assessment required after 4 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 NZ Hospital. and 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 or 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 or Patient has severe chronic localised genital or flexural plaque psoriasis where the plaques or lesions have been present for at least 6 months from the time of initial diagnosis, and with a Dermatology Life Quality Index (DLQI) score greater than 10 and Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin and A PASI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior treatment course, preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course and The most recent PASI or DQLI assessment is no more than 1 month old at the time of application 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, foot, genital or flexural areas, at least 2 of the 3 PASI symptom sub scores for erythema, thickness and scaling are rated as severe or very severe, and for the face, palm of a hand or sole of a foot 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) Patient's PASI score has reduced by 75% or more (PASI 75) as compared to baseline PASI prior to commencing secukinumab or Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing secukinumab or Patient had severe chronic localised genital or flexural plaque psoriasis at the start of treatment and The patient has experienced a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-treatment baseline value or Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing secukinumab and Secukinumab to be administered at a maximum dose of 300 mg monthly

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PRES	CRIB	ER		PATIENT:		
Name	e:			Name:		
Ward	:			NHI:		
Secu	ıkinı	ımab -	- continued			
			ylosing spondylitis, second-line biologic quired after 3 months			
			c boxes where appropriate)			
and		Prescribe Hospital.	ed by, or recommended by a rheumatologist, or in accordance	ce with a protocol or guideline that has been endorsed by the Health NZ		
	and	O The	e patient has had an initial Special Authority approval for ad	alimumab and/or etanercept for ankylosing spondylitis		
		$\bigcap_{\mathbf{or}}$	The patient has experienced intolerable side effects from	n a reasonable trial of adalimumab and/or etanercept		
		С	Following 12 weeks of adalimumab and/or etanercept treated and/or etanercept for ankylosing spondylitis	eatment, the patient did not meet the renewal criteria for adalimumab		
Re-a	ssess <b>equis</b>	Sinuation – ankylosing spondylitis, second-line biologic sessment required after 6 months quisites (tick boxes where appropriate)  Prescribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.				
unu	and		llowing 12 weeks initial treatment of secukinumab treatment seline on a 10 point scale, or by 50%, whichever is less	t, BASDAI has improved by 4 or more points from pre-secukinumab		
	and	O Ph	ysician considers that the patient has benefitted from treatm	nent and that continued treatment is appropriate		
	and	O Sec	cukinumab to be administered at doses no greater than 150	mg monthly		

I confirm that the above details are correct:

Signed: ...... Date: .....

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PRES	PRESCRIBER			PATIENT:
Name	ə:			Name:
Ward	:			NHI:
Seci	ukinu	umal	<b>)</b> - c	tinued
Re-a	assess equis	ment ites (1	requ tick b tibed	e arthritis d after 6 months es where appropriate) , or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
and			$\overline{}$	
		and	$\bigcirc$	atient 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
			or	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	$\circ$	atient has had severe active psoriatic arthritis for six months duration or longer
		and	0	atient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg eekly or a maximum tolerated dose
		and	0	atient 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 dose of up to 20 mg daily (or maximum tolerated doses)
		and		Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen, tender joints
			or	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
		and		
				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
			or	O Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour
			or	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
Re-a	assess	ment	requ	riatic arthritis d after 6 months
Prer	_	,		es where appropriate)
and		Prescr Hospit		, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
		or	0	ollowing 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a inically significant response to treatment in the opinion of the physician
			0	ne patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant esponse to prior secukinumab treatment in the opinion of the treating physician
and O Secukinumab to be administered at doses no greater than 300 mg monthly				numab to be administered at doses no greater than 300 mg monthly