Enquiries to Ministry of Health 0800 855 066

## APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:						
Reg No:	First Names:	First Names:						
Name:	Surname:	Surname:						
Address:	DOB:	Address:						
	Address:							
Fax Number:		Fax Number:						
Secukinumab								
Initial application — severe chronic plaque psoriasis – second-line biologic Applications only from a dermatologist or any relevant practitioner on the recommendation of a dermatologist. Approvals valid for 4 months.  Prerequisites(tick boxes where appropriate)  The patient has had an initial Special Authority approval for adalimumab or etanercept, or has trialled infliximab in a Health NZ Hospital, for severe chronic plaque psoriasis  and  The patient has experienced intolerable side effects from adalimumab, etanercept or infliximab  or  The patient has received insufficient benefit from adalimumab, etanercept or infliximab  and  A Psoriasis Area and Severity Index (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  The most recent PASI or DQLI assessment is no more than 1 month old at the time of application								
Initial application — severe chronic plaque psoriasis – first-line biologic Applications only from a dermatologist or any relevant practitioner on the recommendation of a dermatologist. Approvals valid for 4 months.  Prerequisites(tick boxes where appropriate)								
or Patient has severe chibeen present for at lea	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							
A PASI assessment or Derm treatment course, preferably and The most recent PASI or DC  Note: A treatment course is defined as a mi psoriasis, a PASI score of greater than 10, a recent prior treatment; for severe chronic pla for erythema, thickness and scaling are rate	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  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  The most recent PASI or DQLI assessment is no more than 1 month old at the time of application  ent course is defined as a minimum of 12 weeks of treatment. "Inadequate response" is defined as: for whole body severe chronic plaque SI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most atment; for severe chronic plaque psoriasis of the face, hand. foot, genital or flexural areas, at least 2 of the 3 PASI symptom sub scores nickness 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 e, 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 or treatment.							

I confirm the above details are correct and that in signing this form I understand I may be audited.

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Addre	ss:					D	OB:			Address:	
						A	ddress:				
Fax N	umbei	r:								Fax Number:	
Secu	kinu	ımab	- co	ntinued							
Rene	wal –	– sev	ere cl	hronic p	laque psoriasis	– firs	st and second-line l	biologic			
		•		`	,						
			-		practitioner. App e appropriate)	orovals	s valid for 6 months.				
[											
						re has	nas reduced by 75% or more (PASI 75) as compared to baseline PASI prior to commencing				
			or		cukinumab	ermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to					
					mmencing secul			iex (DEQI) Improv	rement of 5 o	or more, as compared to baseline DEQI prior to	
		or									
			and		Patient had severe chronic localised genital or flexural plaque psoriasis at the start of treatment						
									e in the skin a	area affected, or sustained at this level, as	
				or							
				L			natology Quality of L ng secukinumab	ife Index (DLQI) i	improvement	t of 5 or more, as compared to baseline DLQI	
	and [		Secuk	inumab t	to be administere	ed at a	n maximum dose of 3	300 mg monthly			
Į.											
							ond-line biologic er on the recommend	dation of a rheum	atologist. Ap	oprovals valid for 3 months.	
	Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 3 months. <b>Prerequisites</b> (tick boxes where appropriate)										
	[		The pa	atient has	s had an initial S	pecial	Authority approval f	for adalimumab a	nd/or etanero	cept for ankylosing spondylitis	
	and										
		or	□ ·	The patie	ent has experien	ced int	tolerable side effects	s from a reasonal	ole trial of ad	lalimumab and/or etanercept	
					g 12 weeks of actanercept for ank			ept treatment, the	e patient did r	not meet the renewal criteria for adalimumab	
				4114/01	tanoroopt for ann	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	g openaying				
Rene	Renewal — ankylosing spondylitis – second-line biologic										
Current approval Number (if known):											
Applications only from a rheumatologist or medical practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. <b>Prerequisites</b> (tick boxes where appropriate)											
]	-quioi			WHO!							
	L						of secukinumab trea 0%, whichever is less		as improved	by 4 or more points from pre-secukinumab	
	and	_			•	-			t continued tr	reatment is appropriate	
	and	_					at doses no greater than 300 mg monthly				
	L	`	Jecuk	iiiuiiiad I	o de administêre	u ai d	uses no greater that	ii soo iiig inonini)	/		

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Addre	ss:				DOB:	Address:					
					Address:						
Fax Number:						Fax Number:					
Initia Appl	al application	<b>licatio</b> is only	n — fron	psoriatic arthritis n a rheumatologist. Approva	als valid for 6 months.						
		and		Patient has had an initial Special Authority approval for adalimumab, etanercept or infliximab for psoriatic arthritis							
		a.i.a		Patient has experien	ient 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 [	Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg w or a maximum tolerated dose								
		and			or maximum tolerated doses)	about of at loads 2 g per day or longitorings at a					
	Patient has persistent				t symptoms of poorly controlled and active disease in at least 15 swollen, tender joints t symptoms of poorly controlled and active disease in at least four joints from the following: wrist,						
				elbow, knee, ankle, a	and either shoulder or hip						
		and	or	Patient has a C-reac application	tive protein level greater than 15 mg/L measured no m	nore than one month prior to the date of this					
				Patient has an eleva	ted erythrocyte sedimentation rate (ESR) greater than	25 mm per hour					
			or		neasured as patient is currently receiving prednisone the more than three months	nerapy at a dose of greater than 5 mg per day					
_											
	Renewal — psoriatic arthritis  Current approval Number (if known):										
Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months.  Prerequisites(tick boxes where appropriate)											
	Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician										
		or [		The patient demonstrates a	at least a continuing 30% improvement in active joint c mab treatment in the opinion of the treating physician	ount from baseline and a clinically significant					
	and [	s	ecul	kinumab to be administered	at doses no greater than 300 mg monthly						

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