RS2062 - Etanercept

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Psoriatic arthritis - CONTINUATION	
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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	SCRIE	BER		PATIENT:	
Name	ə:			Name:	
Ward	:			NHI:	
Etan	nerce	ept			
Re-a	assess requis	sment sites († Prescr	requ tick b ibed	ticular course juvenile idiopathic arthritis ired after 6 months oxes where appropriate) by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed th NZ Hospital.	
and		and	0	The patient has had an initial Special Authority approval for adalimumab for polyarticular course juvenile idiopathic arthritis (JIA)	
			or	 O The patient has experienced intolerable side effects from adalimumab O The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for polyarticular course JIA 	
	or				
		and and	0	To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance Patient has had polyarticular course JIA for 6 months duration or longer	
			or	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)	
			or	O Moderate or high disease activity (cJADAS10 score of at least 2.5) after a 3-month trial of methotrexate (at the maximum tolerated dose)	
				O Low disease activity (cJADAS10 score between 1.1 and 2.5) after a 6-month trial of methotrexate	
\subseteq					
Re-a	assess	sment	requ	olyarticular course juvenile idiopathic arthritis ired after 6 months oxes where appropriate)	
and		Prescr by the	ibed Heal	by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed th NZ Hospital.	
	and	į		ment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or rance	
				0	Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline
		or	0	On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline	
			_		

Signed: Date:

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PRESC	RIBE	R	PATIENT:
Name:			Name:
Ward:			
Etane	rcep	t - cont	inued
Re-ass	sessm quisite) Pre	ent reques (tick bescribed	articular course juvenile idiopathic arthritis hired after 6 months boxes where appropriate) by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed
and	by	the Hea	Ith NZ Hospital.
		O	The patient has had an initial Special Authority approval for adalimumab for oligoarticular course juvenile idiopathic arthritis (JIA)
		or	 O The patient has experienced intolerable side effects from adalimumab O The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for oligoarticular course JIA
	or		
	a		To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance Patient has had oligoarticular course JIA for 6 months duration or longer
		and or	O At least 2 active 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 greater than 1.5) with poor prognostic features after a 3-month trial of methotrexate (at the maximum tolerated dose)
		or	O High disease activity (cJADAS10 score greater than 4) after a 6-month trial of methotrexate
Re-ass	sessm	ent requ	bligoarticular course juvenile idiopathic arthritis nired after 6 months boxes where appropriate)
and			by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed Ith NZ Hospital.

Subsidised as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolera	nce

Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baselinee

On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline

and

or

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PRE	SCRIE	BER		PATIENT:
Nam	e:			Name:
Ward	d:			NHI:
Etar	nerce	ept - c	onti	ued
Re-a	asses: r equis	sment re sites (tio	equi ck b	s - rheumatoid ed after 6 months xes where appropriate) y, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
and	l	Hospita		y, or recommended by a medinalologist, or in accordance with a protocor or guideline that has been endorsed by the median we
		and	C	The patient has had an initial Special Authority approval for adalimumab for rheumatoid arthritis
			or	O The patient has experienced intolerable side effects
				O The patient has received insufficient benefit to meet the renewal criteria for rheumatoid arthritis
	or			
		and and and and and and	0 0 0 0 0 0 0 0 0	Patient has had rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance Patient has tried and not responded to at least three months of methotrexate at a maximum tolerated dose (unless contraindicated) Patient has tried and not responded to at least three months of methotrexate in combination with sulfasalazine and hydroxychloroquine sulphate at maximum tolerated doses (unless contraindicated) Patient has tried and not responded to at least three months of methotrexate in combination with the maximum tolerated dose of ciclosporin Patient has tried and not responded to at least three months of therapy at the maximum tolerated dose of leflunomide alone or in combination with methotrexate Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen joints 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
Re-a		sment re sites (tio Prescrib NZ Hos O Tr	equi ck b ped pita	thritis - rheumatoid ed after 2 years exes where appropriate) by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health ment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or ance
	and)	Following initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant

Ο	On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count from
	baseline and a clinically significant response to treatment in the opinion of the physician

O Etanercept to be administered at doses no greater than 50 mg every 7 days

response to treatment in the opinion of the physician

I confirm that the above details are correct:

or

and

Signed: Date:

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PRESC	RIBE	ER					PATIENT:
Name:							Name:
Ward: .							NHI:
Etaner	rcep	ot - a	conti	nued			
					spondylitis		
			•		fter 6 months	to)	
Prereq	uisi	les (l		oxes	where appropria	le)	
		rescri ospita		by, or	recommended	by a rheumatologist, o	or in accordance with a protocol or guideline that has been endorsed by the Health NZ
and		ospila	aı.				
		and	О	The	patient has had	an initial Special Autho	nority approval for adalimumab for ankylosing spondylitis
		unu	or	0	The patient ha	s experienced intolera	able side effects from adalimumab
				0	The patient ha		t benefit from adalimumab to meet the renewal criteria for adalimumab for
o	or (
	ſ		Ο	Patie	nt has a confirm	ed diagnosis of ankyl	rlosing spondylitis present for more than six months
		and	$\overline{\mathbf{O}}$				
		and	\mathbf{O}	Patie	nt has low back	pain and stiffness tha	at is relieved by exercise but not by rest
		anu	О	Patie	nt has bilateral	sacroiliitis demonstrate	ted by plain radiographs. CT or MRI scan
		and					
			\bigcirc	drug	s (NSAIDs), in c	spondylitis has not res ombination with anti-u ankylosing spondylitis	ulcer therapy if indicated, while patient was undergoing at least 3 months of a regular
		and	_	CACIO			•
				0	Bath Ankylosin	g Spondylitis Metrolog	bgy Index (BASMI) measures: a modified Schober's test of less than or equal to
			or	Ο	Patient has lim gender (see N		sion by at least 2.5 cm below the average normal values corrected for age and
		and			gender (see h	5103)	
		anu	О	Bath	Ankylosing Spo	ndylitis Disease Activi	vity Index (BASDAI) of at least 6 on a 0-10 scale
					, , ,		
						for age and gender:	Name: Name: NHI: NHI: umatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Special Authority approval for adalimumab for ankylosing spondylitis anced intolerable side effects from adalimumab ad insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for nosis of ankylosing spondylitis present for more than six months d stiffness that is relieved by exercise but not by rest is demonstrated by plain radiographs, CT or MRI scan tis has not responded adequately to treatment with two or more non-steroidal anti-inflammatory for with anti-lucer therapy if indicated, while patient was undergoing at least 3 months of a regular ing spondylitis frontion of the lumbar spine in the sagittal and the frontal planes as determined by the following lybitis Metrology Index (BASMI) measures: a modified Schober's test of less than or equal to least more and lett and right) f chest expansion by at least 2.5 cm below the average normal values corrected for age and Disease Activity Index (BASDAI) of at least 6 on a 0-10 scale the completion of the 3 month exercise trial, but prior to ceasing NSAID treatment. The BASDAI and gender: nale om cm om cm om cm
			Age		Male	Female	
			18-2		7.0 cm	5.5 cm	
			25-3	4	7.5 cm	5.5 cm	
			35-4	4	6.5 cm	4.5 cm	
			45-5	4	6.0 cm	5.0 cm	
			55-6	4	5.5 cm	4.0 cm	
			65-7	4	4.0 cm	4.0 cm	

75+

3.0 cm

Signed: Date:

2.5 cm

PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:
Etanercept - continued	
and Hospital.	nent and that continued treatment is appropriate
INITIATION – psoriatic arthritis Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a rheumatologist, or in accordar Hospital.	nce with a protocol or guideline that has been endorsed by the Health NZ
O The patient has had an initial Special Authority approva and O The patient has experienced intolerable side effect or O The patient has received insufficient benefit from adalimumab or secukinumab for psoriatic arthritis	cts from adalimumab or secukinumab adalimumab or secukinumab to meet the renewal criteria for
weekly or a maximum tolerated dose and Patient has tried and not responded to at least three mo a dose of up to 20 mg daily (or maximum tolerated dose and O Patient has persistent symptoms of poorly control	onths of oral or parenteral methotrexate at a dose of at least 20 mg
or O Patient has an elevated erythrocyte sedimentation	n 15 mg/L measured no more than one month prior to the date of this n rate (ESR) greater than 25 mm per hour dy receiving prednisone therapy at a dose of greater than 5 mg per day

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PRES	CRIB	ER			PATIENT:
Name	:				Name:
Ward:					NHI:
Etan	erce	pt -	- conti	nued	
Re-a	ssess	men	t requi	soriatic arthritis red after 6 months oxes where appropriate)	
(and		resc losp		by, or recommended by a rheumatologist, or in accordance	ce with a protocol or guideline that has been endorsed by the Health NZ
		or	0 0	clinically significant response to treatment in the opinion	rovement in active joint count from baseline and a clinically significant
	and (С	Etane	ercept to be administered at doses no greater than 50 mg	every 7 days
Re-a	ssess equisi	men i tes	t requi (tick b cribed	chronic plaque psoriasis, prior TNF use red after 4 months oxes where appropriate) by, or recommended by a dermatologist, or in accordance	e with a protocol or guideline that has been endorsed by the Health NZ
	(and	С	The p	patient has had an initial Special Authority approval for ad	alimumab for severe chronic plaque psoriasis
		or	0 0	The patient has experienced intolerable side effects from The patient has received insufficient benefit from adalim- plaque psoriasis	n adalimumab umab to meet the renewal criteria for adalimumab for severe chronic
	and (C	Patie	nt must be reassessed for continuation after 3 doses	

I confirm that the above details are correct:

Signed: Date:

PRE	SCRIE	BER			PATIENT:
Name	e:				Name:
Ward	I:				NHI:
Etar	nerce	ept	- conti	nued	
				e chronic plaque psoriasis, treatment-naive ired after 4 months	
				oxes where appropriate)	
and		Preso Hosp		by, or recommended by a dermatologist, or in accordance	e with a protocol or guideline that has been endorsed by the Health NZ
		or	0	10, where lesions have been present for at least 6 month Patient has severe chronic plaque psoriasis of the face, been present for at least 6 months from the time of initia	or palm of a hand or sole of a foot, where the plaque or plaques have
	and and	0	follow	least 6 months from the time of initial diagnosis, and with nt has tried, but had an inadequate response (see Note) ving (at maximum tolerated doses unless contraindicated) SI assessment or Dermatology Quality of Life Index (DLC	n a Dermatology Life Quality Index (DLQI) score greater than 10 to, or has experienced intolerable side effects from, at least three of the
Note	and	0	cessa The r	ation of each prior treatment course most recent PASI or DLQI assessment is no more than 1	
while face seve	e still (, hanc ere, ar	on tre d, foo nd for	eatmei t, geni the fa	nt but no longer than 1 month following cessation of the r tal or flexural areas at least 2 of the 3 PASI symptom sul	nost recent prior treatment; for severe chronic plaque psoriasis of the oscores for erythema, thickness and scaling are rated as severe or very d is 30% or more of the face, palm of a hand or sole of a foot, as assessed

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PRESCRIBER	PATIENT:	
Name:	Name:	
Ward:	NHI:	

Etanercept - continued

CONTINUATION – severe chronic plaque psoriasis Re-assessment required after 6 months **Prerequisites** (tick boxes where appropriate)

	and	O Patient had "whole body" severe chronic plaque psoriasis at the start of treatment
		 Following each prior etanercept treatment course the patient has a PASI score which is reduced by 75% or more, or is sustained at this level, when compared with the pre-etanercept treatment baseline value Following each prior etanercept treatment course the patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, when compared with the pre-treatment baseline value
or		
	and	O Patient had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment
		O Following each prior etanercept treatment course the patient has a reduction in the PASI symptom subscores for all 3 of erythema, thickness and scaling, to slight or better, or sustained at this level, as compared to the treatment course baseline values
		O Following each prior etanercept treatment course the patient has a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-etanercept treatment baseline value
or		
	and	O Patient had severe chronic localised genital or flexural plaque psoriasis at the start of treatment
		O 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
		O Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing etanercept
and	Etaner	rcept to be administered at doses no greater than 50 mg every 7 days
		ma gangrenosum
requisites	(tick bo	ixes where appropriate)
O Pres Hosp		by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
O	Patien	t has pyoderma gangrenosum*
and		t has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclospori oprine, or methotrexate) and not received an adequate response

O A maximum of 8 doses

Note: Indications marked with * are unapproved indications.

PRESCRIBER							PATIENT:	
Name:							Name:	
Ward:							NHI:	
Etan	erce	pt-a	conti	led				
CONTINUATION – pyoderma gangrenosum Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the H Hospital.								
and	 Patient has shown clinical improvement and Patient continues to require treatment and A maximum of 8 doses 							
INITIATION – adult-onset Still's disease Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital. and								
		and	or	The patieThe patieThe patieThe patie	ent has been start ent has experienc ent has received i	ted on tocilizumab for A ed intolerable side effect	oproval for etanercept for adult-onset Still's disease (AOSD) OSD in a Health NZ Hospital ets from etanercept and/or tocilizumab at least a three-month trial of adalimumab and/or tocilizumab such that	
	or	and (and	С С С	Patient has trie Intiinflammator	d and not respon ry drugs (NSAIDs	ded to at least 6 month and methotrexate	chi criteria (J Rheumatol 1992;19:424-430) s of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal atrolled and active disease	
CONTINUATION – adult-onset Still's disease Re-assessment required after 6 months Prerequisites (tick box where appropriate)								
and	н С	lospita	al.			atologist, or in accordar inflammatory markers a	and functional status	

PRES	SCRIE	BER		PATIENT:				
Name	e:			Name:				
Ward	:			NHI:				
Etan	erce	ept -	cont	inued				
Re-a	ssess equis	ites	t requ (tick b ribed	erentiated spondyloarthritis hired after 6 months boxes where appropriate) by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ				
and	and	0 0	Patie wrist Patie	ent has undifferentiated peripheral spondyloarthritis* with active peripheral joint arthritis in at least four joints from the following: , elbow, knee, ankle, and either shoulder or hip ent 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 mum tolerated dose				
	and (and	0		ent has tried and not responded to at least three months of sulfasalazine at a dose of at least 2 g per day (or maximum tolerated				
	and	0	Patie	ent has tried and not responded to at least three months of leflunomide at a dose of up to 20 mg daily (or maximum tolerated dose)				
			0	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	0	Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour measured no more than one month prior to the date of this application				
		or	0	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				
Note	: Indi	catio	ns ma	arked with * are unapproved indications.				
Re-a	ssess	men	t requ	undifferentiated spondyloarthritis hired after 6 months boxes where appropriate)				
		or	0	Applicant is a rheumatologist				
			0	Applicant is a Practitioner and confirms that a rheumatologist has provided a letter, email or fax recommending that the patient continues with etanercept treatment				
	and	or	0	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				
			Ο	The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior etanercept treatment in the opinion of the treating physician				
and O Etanercept to be administered at doses no greater than 50 mg dose every 7 days								

Signed: Da	ate:
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