## RS2062 - Etanercept

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PRES	SCRIE	BER	PATIENT:
Name	ə:		Name:
Ward	:		NHI:
Etan	erce	ept	
Re-a	equis	sment sites ( Presc	olyarticular course juvenile idiopathic arthritis required after 6 months tick boxes where appropriate) ribed by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed Health NZ Hospital.
		and	The patient has had an initial Special Authority approval for adalimumab for polyarticular course juvenile idiopathic arthritis (JIA)
			Or The patient has experienced intolerable side effects from adalimumab  The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for polyarticular course JIA
	or	and	O Patient has had polyarticular course JIA for 6 months duration or longer
Re-a	assess	sment	N – polyarticular course juvenile idiopathic arthritis required after 6 months tick boxes where appropriate)
and			ribed by, or recommended by a rheumatologist or named specialist, or in accordance with a protocol or guideline that has been endorsed Health NZ Hospital.
	and		Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance
		or	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  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

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PRE	SCRIE	BER		PATIENT:
Name	e:			Name:
Ward	l:			NHI:
Etar	erce	ept -	conti	nued
Re-a	assess requis	sment <b>sites</b> ( Presci	requ tick b ribed	rticular 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 oligoarticular course juvenile idiopathic arthritis (JIA)
		and	or	The patient has experienced intolerable side effects from adalimumab     The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for oligoarticular course JIA
	or	and	$\circ$	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  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)  O High disease activity (cJADAS10 score greater than 4) after a 6-month trial of methotrexate
Re-a	assess requis	sment <b>sites</b> ( Presci	requ tick b ribed	ligoarticular 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		Subs	idised as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance
		or	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 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

I confirm that the above details are correct:

I confirm that the above details are correct:

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

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PRES	CRIB	BER			PATIENT:
Name:	:				Name:
Ward:					NHI:
Etane	erce	pt -	conti	nued	
				s - rheumatoid	
				red after 6 months  expectations are series of the series	
(	`				de constate de la constante de
		rescr Hospit		by, or recommended by a	rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
and			$\overline{\bigcirc}$	The constant has been been as to	
		and		The patient has had an ir	nitial Special Authority approval for adalimumab for rheumatoid arthritis
			۵.	O The patient has exp	perienced intolerable side effects
			or	O The patient has red	ceived insufficient benefit to meet the renewal criteria for rheumatoid arthritis
	or				
			O		oid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP)
		and		antibody positive) for six	months duration or longer
			$\bigcirc$	Treatment is to be used a or intolerance	as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity
		and	$\bigcirc$		responded to at least three months of methotrexate at a maximum tolerated dose (unless contraindicated)
		and	$\bigcirc$		
					responded to at least three months of methotrexate in combination with sulfasalazine and hydroxychloroquinerated doses (unless contraindicated)
		and		O Patient has tried an	nd not responded to at least three months of methotrexate in combination with the maximum tolerated
			or	dose of ciclosporin	in not responded to at least time months of methodexate in combination with the maximum tolerated
					nd not responded to at least three months of therapy at the maximum tolerated dose of leflunomide ation with methotrexate
		and		aione or in combina	anon with methodexate
				O Patient has persiste	ent symptoms of poorly controlled and active disease in at least 15 swollen joints
			or	O Patient has persiste	ent symptoms of poorly controlled and active disease in at least four joints from the following: wrist,
					, and either shoulder or hip
				rthritis - rheumatoid red after 2 years	
				exes where appropriate)	
( and		Prescr NZ Ho			ny relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health
	and			nent is to be used as an a	adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or
		or	0		t, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant the opinion of the physician
			0		tions, the patient demonstrates at least a continuing 30% improvement in active joint count from significant response to treatment in the opinion of the physician
	and	$\bigcirc$	E+o∽-	roopt to be administered	at desce no greater than 50 mg every 7 deve
			Liane	rcept to be administered	at doses no greater than 50 mg every 7 days
			_		

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RESCRIBER			PATIENT:
me:			Name:
ard:			NHI:
anercept	- continued	1	
e-assessmer rerequisites	nt required (tick boxes cribed by, c	g spondylitis after 6 months where appropria or recommended	ate) I by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
an		patient has had	an initial Special Authority approval for adalimumab for ankylosing spondylitis
	or O	The patient ha	as experienced intolerable side effects from adalimumab
	ů O	The patient ha ankylosing spo	as received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for ondylitis
or			
an an an	Pati	ient has low back ient has bilateral ient's ankylosing gs (NSAIDs), in c rcise regimen for Patient has lim Bath Ankylosir 4 cm and lumb Patient has lim gender (see N	k pain and stiffness that is relieved by exercise but not by rest sacroiliitis demonstrated by plain radiographs, CT or MRI scan spondylitis has not responded adequately to treatment with two or more non-steroidal anti-inflammatory combination with anti-ulcer therapy if indicated, while patient was undergoing at least 3 months of a regular rankylosing spondylitis mitation of motion of the lumbar spine in the sagittal and the frontal planes as determined by the following ng Spondylitis Metrology Index (BASMI) measures: a modified Schober's test of less than or equal to bar side flexion measurement of less than or equal to 10 cm (mean of left and right) mitation of chest expansion by at least 2.5 cm below the average normal values corrected for age and lotes) ondylitis Disease Activity Index (BASDAI) of at least 6 on a 0-10 scale
			mined at the completion of the 3 month exercise trial, but prior to ceasing NSAID treatment. The BASDAI
easure must	be no more	e than 1 month o	old at the time of starting treatment. d for age and gender: Female 5.5 cm 5.5 cm 4.5 cm 5.0 cm 4.0 cm 4.0 cm 2.5 cm

I confirm that the above details are correct:

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PRESCRIBER	PATIENT:
Name:	
Ward:	NHI:
Etanercept - contin	nued
Re-assessment requi	nkylosing spondylitis red after 6 months oxes where appropriate) oy, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
Follow points and Physic and	wing 12 weeks' initial treatment and for subsequent renewals, treatment has resulted in an improvement in BASDAI of 4 or more of from pre-treatment baseline on a 10 point scale, or an improvement in BASDAI of 50%, whichever is less can considers that the patient has benefited from treatment and that continued treatment is appropriate procept to be administered at doses no greater than 50 mg every 7 days
and	The patient has had an initial Special Authority approval for adalimumab or secukinumab for psoriatic arthritis  The patient has experienced intolerable side effects from adalimumab or secukinumab  The patient has received insufficient benefit from adalimumab or secukinumab to meet the renewal criteria for adalimumab or secukinumab for psoriatic arthritis
and on the same of	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  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  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  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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PRES	CRIE	BER			PATIENT:
Name	e:				Name:
Ward:	:				NHI:
Etan	erce	pt -	- conti	inued	
Re-a	ssess	smen	t requ	osoriatic arthritis ired after 6 months oxes where appropriate)	
and		Preso Hosp		by, or recommended by a rheumatologist, or in accordance	ee with a protocol or guideline that has been endorsed by the Health NZ
		or	O O	clinically significant response to treatment in the opinion	ovement in active joint count from baseline and a clinically significant
	and	0	Etane	ercept to be administered at doses no greater than 50 mg	
Re-a	ssess	smen	t requ	e chronic plaque psoriasis, prior TNF use ired after 4 months poxes where appropriate)	
( and		Preso Hosp		by, or recommended by a dermatologist, or in accordance	with a protocol or guideline that has been endorsed by the Health NZ
	and	0	The p	patient has had an initial Special Authority approval for add	alimumab for severe chronic plaque psoriasis
	and.	or	O O	The patient has experienced intolerable side effects from The patient has received insufficient benefit from adalimuplaque psoriasis	adalimumab umab to meet the renewal criteria for adalimumab for severe chronic
	and	0	Patie	nt must be reassessed for continuation after 3 doses	

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ESCRIBER		PATIENT:
me:		Name:
ard:		NHI:
anercept	- continued	
	severe chronic plaque psoriasis, treatment-naive t required after 4 months	
erequisites	(tick boxes where appropriate)	
Preso Hosp	,	nce with a protocol or guideline that has been endorsed by the Health NZ
or	O Patient has "whole body" severe chronic plaque psoria 10, where lesions have been present for at least 6 mor	asis with a Psoriasis Area and Severity Index (PASI) score of greater than on the time of initial diagnosis
or	O Patient has severe chronic plaque psoriasis of the face been present for at least 6 months from the time of init	e, or palm of a hand or sole of a foot, where the plaque or plaques have ial diagnosis
		plaque psoriasis where the plaques or lesions have been present for at rith a Dermatology Life Quality Index (DLQI) score greater than 10
and	Patient has tried, but had an inadequate response (see Note following (at maximum tolerated doses unless contraindicate	e) to, or has experienced intolerable side effects from, at least three of the ed): phototherapy, methotrexate, ciclosporin, or acitretin
and		QI) assessment has been completed for at least the most recent prior preferably while still on treatment but no longer than 1 month following
	The most recent PASI or DLQI assessment is no more than	1 month old at the time of initiation
nile still on tre ce, hand, foo evere, and for	eatment but no longer than 1 month following cessation of the t, genital or flexural areas at least 2 of the 3 PASI symptom s	plaque psoriasis, a PASI score of greater than 10, as assessed preferably a most recent prior treatment; for severe chronic plaque psoriasis of the subscores for erythema, thickness and scaling are rated as severe or very ed is 30% or more of the face, palm of a hand or sole of a foot, as assessedation of the most recent prior treatment.

RESCRIBER	PATIENT:
ame:	Name:
/ard:	NHI:
tanercept - continued	
CONTINUATION – severe chror Re-assessment required after 6 r Prerequisites (tick boxes where	nonths
O Patie and	nt had "whole body" severe chronic plaque psoriasis at the start of treatment
or O	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 Patie	nt had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment
or 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  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 Patie	nt had severe chronic localised genital or flexural plaque psoriasis at the start of treatment
or 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
	Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing etanercept
and Etanercept to be	administered at doses no greater than 50 mg every 7 days
IITIATION – pyoderma gangre	nosum
rerequisites (tick boxes where	
Prescribed by, or recome Hospital.	mended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
O Patient has pyode	erma gangrenosum*
azathioprine, or n	red three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclosporin, nethotrexate) and not received an adequate response
O A maximum of 8	
Note: Indications marked with * a	are unapproved indications.

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PRES	CRIB	ER			PATIENT:
Name	:				Name:
Ward:					NHI:
Etan	erce	pt - ۵	onti	nued	
	iaiupe P	rescritospita	ck b bed al.	oxes	ma gangrenosum where appropriate) recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ shown clinical improvement
	and ( and	$\sim$			tinues to require treatment of 8 doses
Re-a	ssessi <b>equisi</b> P	ment i <b>tes</b> (t	equick b	red a	Still's disease ter 6 months where appropriate) recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
		and	or	0	The patient has had an initial Special Authority approval for etanercept for adult-onset Still's disease (AOSD)  The patient has been started on tocilizumab for AOSD in a Health NZ Hospital  The patient has experienced intolerable side effects from etanercept and/or tocilizumab  The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or tocilizumab such that they do not meet the renewal criteria for AOSD
	or	and	) ) )	Patie antii	Int diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 1992;19:424-430)  Int has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal flammatory drugs (NSAIDs) and methotrexate  Int has persistent symptoms of disabling poorly controlled and active disease
Re-a	ssessi equisi P H	ment i <b>tes</b> (t Prescri lospita	equick b bed al.	red a ox wh by, or	nset Still's disease ter 6 months ere appropriate) recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
	<b>У</b> т	he pa	tient	has	sustained improvement in inflammatory markers and functional status

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SCRIB	ER		PATIENT:
e:			
d:			NHI:
nerce	pt -	conti	nued
TIATION assessi requisi	N - u mentites (	ribed tal.  Patie wrist,  Patie maxii  Patie dose	erentiated spondyloarthritis ired after 6 months oxes where appropriate)  by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ  int 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  int 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  int 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
e: Indic	or		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  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  arked with * are unapproved indications.
e. maic	Jalioi	15 1116	Thed with are unapproved indications.
assessi	ment	t requ	Indifferentiated spondyloarthritis ired after 6 months inoxes where appropriate)
0.1	or	O O	Applicant is a rheumatologist  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	O O	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  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
	$\overline{}$		

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