SA2399 - Etanercept

Arthritis - rheumatoid - Renewal	10
Arthritis - rheumatoid - Initial application	
Adult-onset Still's disease - Initial application	
Adult-onset Still's disease - Renewal	2
Ankylosing spondylitis - Initial application	
Ankylosing spondylitis - Renewal	
Oligoarticular course juvenile idiopathic arthritis - Initial application	
Oligoarticular course juvenile idiopathic arthritis - Renewal	
Polyarticular course juvenile idiopathic arthritis - Initial application	
Polyarticular course juvenile idiopathic arthritis - Renewal	5
Psoriatic arthritis - Initial application	7
Psoriatic arthritis - Renewal	8
Pyoderma gangrenosum - Initial application	
Pyoderma gangrenosum - Renewal	8
Severe chronic plaque psoriasis - Initial application	11
Severe chronic plaque psoriasis - Renewal	12
Undifferentiated spondyloarthritis - Initial application	
Undifferentiated spondyloarthritis - Renewal	
Chambroniated openational frontieral minimum m	

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)			p or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg I	No:			First Names:	First Names:
Name	e:			Surname:	Surname:
Addre	ess:			DOB:	Address:
				Address:	
Fax N	lumber	:			Fax Number:
Etan	erce	pt			
App	lication	ns only	n — adult-onset Still's disease from a rheumatologist. Approva (a boxes where appropriate)	ls valid for 6 months.	
		and	or	n initial Special Authority approval for adalimumab fo started on tocilizumab for AOSD in a Health NZ Hosp	
			or The patient has received	ienced intolerable side effects from adalimumab and/ ved insufficient benefit from at least a three-month tria renewal criteria for AOSD	
Patient diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 19 and Patient has tried and not responded to at least 6 months of glucocorticosteroids at anti-inflammatory drugs (NSAIDs) and methotrexate				·	
		and [Patient has persistent symp	toms of disabling poorly controlled and active disease	
Ren	ewal –	– adult	-onset Still's disease		
Appl	ication	s only f	Number (if known): from a rheumatologist or Practition toxes where appropriate)	oner on the recommendation of a rheumatologist. Ap	provals valid for 6 months.
		or [Applicant is a rheumatologis Applicant is a Practitioner are continues with etanercept trees.	nd confirms that a rheumatologist has provided a lette	er, email or fax recommending that the patient
	and	Tr	ne patient has a sustained impro	vement in inflammatory markers and functional status	S

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APPLICAI	NT (stamp o	or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:			. First Names:	First Names:
Name:			Surname:	Surname:
Address:			. DOB:	Address:
			. Address:	
Fax Numb	er:			Fax Number:
Etanerc	ept - contir	nued		
Applicati	ons only from	ankylosing spondylitis m a rheumatologist. Approposes where appropriate)	vals valid for 6 months.	
	and	The patient has exp	tial Special Authority approval for adalimumab f erienced intolerable side effects from adalimum eived insufficient benefit from adalimumab to me	
or		-17		
	and and	Patient has low back pain	agnosis of ankylosing spondylitis present for mo and stiffness that is relieved by exercise but not liitis demonstrated by plain radiographs, CT or	t by rest
	and		nation with anti-ulcer therapy if indicated, while	nt with two or more non-steroidal anti-inflammatory patient was undergoing at least 3 months of a regular
	or	Bath Ankylosing Sprand lumbar side flex Patient has limitation	ondylitis Metrology Index (BASMI) measures: a ion measurement of less than or equal to 10 cm	nd the frontal planes as determined by the following modified Schober's test of less than or equal to 4 cm in (mean of left and right) he average normal values corrected for age and
	and	gender (see Notes) A Bath Ankylosing Spondy	viitis Disease Activity Index (BASDAI) of at least	t 6 on a 0-10 scale
measure Average i 18-24 yea 25-34 yea 35-44 yea 45-54 yea 55-64 yea 65-74 yea	must be no normal ches ars - Male: 7 ars - Male: 6 ars - Male: 6 ars - Male: 5 ars - Male: 5		ne time of initial application.	but prior to ceasing NSAID treatment. The BASDAI

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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:
Etanercept - continued		
Renewal — ankylosing spondylitis		
Current approval Number (if known):		
, ,	ractitioner on the recommendation of a rheumatologist. Ap	oprovals valid for 6 months.
Prerequisites(tick boxes where appropriate)		
or	ologist	
Applicant is a Practitio continues with etanero	ner and confirms that a rheumatologist has provided a lette cept treatment	er, email or fax recommending that the patient
and	·	
	eatment and for subsequent renewals, treatment has resul- seline on a 10 point scale, or an improvement in BASDAI o	
and		
and and	patient has benefited from treatment and that continued tre	eatment is appropriate
Etanercept to be administered	ed at doses no greater than 50 mg every 7 days	
Initial application — polyarticular course	iuvanila idionathic arthritic	
Applications only from a named specialist of	r rheumatologist. Approvals valid for 6 months.	
Prerequisites(tick boxes where appropriate)		
The patient has had a	n initial Special Authority approval for adalimumab for polya	articular course juvenile idiopathic arthritis (JIA)
and		
or The patient has	experienced intolerable side effects from adalimumab	
The patient has course JIA	received insufficient benefit from adalimumab to meet the	renewal criteria for adalimumab for polyarticular
or		
	nct to methotrexate therapy or monotherapy where use of i	methotrexate is limited by toxicity or intolerance
and		
and	ticular course JIA for 6 months duration or longer	
	e joints and at least 3 joints with limited range of motion, pa	ain or tenderness after a 3-month trial of
or	t the maximum tolerated dose)	
tolerated dose)	h disease activity (cJADAS10 score of at least 2.5) after a	ડ-montn trial of methotrexate (at the maximum
or Low disease act	tivity (cJADAS10 score between 1.1 and 2.5) after a 6-mon	th trial of methotrexate

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APPLICANT (stamp or sticker acceptable)			r sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:				First Names:	First Names:
Name: .				Surname:	Surname:
Address:	:			DOB:	Address:
				Address:	
Fax Num					Fax Number:
			cular course juvenile idiopa	nthic arthritis	
Application	tions only 6 months	from	nber (if known): n a named specialist, rheuma xes where appropriate)	tologist or Practitioner on the recommendation of a n	named specialist or rheumatologist. Approvals
a	S	ubsi	dised as an adjunct to metho	trexate therapy or monotherapy where use of methot	rexate is limited by toxicity or intolerance
			physician's global assessme On subsequent reapplicatior continued improvement in ph oligoarticular course juver	ns, the patient demonstrates at least a continuing 309 hysician's global assessment from baseline hile idiopathic arthritis	
	-		n a named specialist or rheur xes where appropriate)	matologist. Approvals valid for 6 months.	
	and		The patient has had an initia	I Special Authority approval for adalimumab for oligo	articular course juvenile idiopathic arthritis (JIA)
			The patient has exper	ienced intolerable side effects from adalimumab	
		or	The patient has receive course JIA	red insufficient benefit from adalimumab to meet the I	renewal criteria for adalimumab for oligoarticular
o	r	_			
	and		To be used as an adjunct to	methotrexate therapy or monotherapy where use of r	methotrexate is limited by toxicity or intolerance
	and		Patient has had oligoarticula	r course JIA for 6 months duration or longer	
		or	At least 2 active joints maximum tolerated do	s with limited range of motion, pain or tenderness after use)	er a 3-month trial of methotrexate (at the
		or		ase activity (cJADAS10 score greater than 1.5) with $\mbox{\sc parameter}$ naximum tolerated dose)	poor prognostic features after a 3-month trial of
		J.	High disease activity (cJADAS10 score greater than 4) after a 6-month trial	of methotrexate

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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:				
Etanercept - continued						
Renewal — oligoarticular course juvenile idiop	athic arthritis					
Current approval Number (if known):						
Applications only from a named specialist, rheuma valid for 6 months.	tologist or Practitioner on the recommendation of a n	amed specialist or rheumatologist. Approvals				
Prerequisites(tick boxes where appropriate)						
Subsidised as an adjunct to metho	Subsidised as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance					
Following 3 to 4 months' init physician's global assessme	e in active joint count and an improvement in					
On subsequent reapplication	ns, the patient demonstrates at least a continuing 30% hysician's global assessment from baseline	6 improvement in active joint count and				

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)			stick	er acceptable)	PATIENT NHI:	REFERRER Reg No:	
Reg N	Reg No:					First Names:	First Names:
Name	:					Surname:	Surname:
Addre	ss:					DOB:	Address:
						Address:	
_		r: pt - <i>c</i>					Fax Number:
App	icatio	ns only	or	The p Patie Patie Pratie	The patient has exper The patient has received or secukinumab for patient has had severe active that has tried and not respect to the patient has persistent elbow, knee, ankle, are patient has a C-reactive application	al Special Authority approval for adalimumab or secukienced intolerable side effects from adalimumab or secukinum and insufficient benefit from adalimumab or secukinum coriatic arthritis The psoriatic arthritis for six months duration or longer ponded to at least three months of oral or parenteral	methotrexate at a dose of at least 20 mg weekly dose of at least 2 g per day or leflunomide at a at least 15 swollen, tender joints at least four joints from the following: wrist,
						asured as patient is currently receiving prednisone theore than three months	erapy at a dose of greater than 5 mg per day

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APPL	ICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:				
Reg N	0:	First Names:	First Names:				
Name		Surname:	Surname:				
Addre	ss:	DOB:	Address:				
		Address:					
Fax N	umber:		Fax Number:				
Etan	ercept - continued						
Rene	wal — psoriatic arthritis						
Curre	ent approval Number (if known):						
	, ,	oner on the recommendation of a rheumatologist. Ap	provals valid for 6 months.				
Prere	equisites(tick boxes where appropriate)						
	Applicant is a rheumatologis	†					
	or						
	continues with etanercept tre	nd confirms that a rheumatologist has provided a lette eatment	er, email or tax recommending that the patient				
	and						
		tial treatment, the patient has at least a 50% decreas e to treatment in the opinion of the physician	e in active joint count from baseline and a				
	or		ount from baseline and a dinically significant				
	response to prior etanercept	least a continuing 30% improvement in active joint or treatment in the opinion of the treating physician	burit from baseline and a clinically significant				
	and						
	Etanercept to be administered at d	oses no greater than 50 mg every 7 days					
Initia	I application — pyoderma gangrenosum						
	ications only from a dermatologist. Approvals equisites (tick boxes where appropriate)	valid for 4 months.					
]	The state of the s						
	Patient has pyoderma gangrenosu	m*					
	Patient has received three months	of conventional therapy including a minimum of three	e pharmaceuticals (e.g. prednisone, ciclosporine,				
	azathioprine, or methotrexate) and not received an adequate response and						
	A maximum of 8 doses						
Note: Indications marked with * are unapproved indications.							
Renewal — pyoderma gangrenosum							
Current approval Number (if known):							
Applications only from a dermatologist or Practitioner on the recommendation of a dermatologist. Approvals valid for 4 months.							
Prere	Prerequisites(tick boxes where appropriate)						
	Patient has shown clinical improve	ment					
	Patient continues to require treatm	ent					
	and						
	A maximum of 8 doses						

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APPLICANT (stamp or sticker acceptable)			mp o	sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:					First Names:	First Names:
Name:					Surname:	Surname:
Addre	Address:				DOB:	Address:
					Address:	
Fax N	Numbe	 er:				Fax Number:
Etan	erce	pt - c	contir	ued		
App	licatio	ns onl	y fror	The patient has exper	al Special Authority approval for adalimumab for rheurienced intolerable side effects yed insufficient benefit to meet the renewal criteria for	
		and and and		antibody positive) for six moderance Treatment is to be used as a intolerance Patient has tried and not resulphate (at maximum tolerance)	an adjunct to methotrexate therapy or monotherapy will sponded to at least three months of methotrexate at a	here use of methotrexate is limited by toxicity or maximum tolerated dose (unless contraindicated) ombination with sulfasalazine and hydroxychloroquine
		and		alone or in combination Patient has persistent Patient has persistent	not responded to at least three months of therapy at to me with methotrexate symptoms of poorly controlled and active disease in symptoms of poorly controlled and active disease in a deither shoulder or hip	at least 15 swollen joints
	1					

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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:		
Etanercept - continued				
Current approval Number (if known):				
or response to treatment in the	e patient has at least a 50% decrease in active joint opinion of the physician as, the patient demonstrates at least a continuing 30% ificant response to treatment in the opinion of the physician	6 improvement in active joint count from		
and Etanercept to be administered at d	oses no greater than 50 mg every 7 days			

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APPLICANT (stamp or sticker acceptable)				er acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:					First Names:	First Names:
Name:					Surname:	Surname:
Address:					DOB:	Address:
					Address:	
	er:					Fax Number:
Initial app	olication ons only	fron	sevei n a de xes w	patient has had an initia	vant practitioner on the recommendation of a dermate all Special Authority approval for adalimumab for sever ienced intolerable side effects from adalimumab ved insufficient benefit from adalimumab to meet the recommendation of a dermate all special Authority approval for adalimumab for sever increase.	re chronic plaque psoriasis
Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) segreter 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 have been present for at least 6 months from the time of initial diagnosis. Patient has severe chronic localised genital or flexural plaque psoriasis where the plaques or lesions have been to for at least 6 months from the time of initial diagnosis, and with a Dermatology Life Quality Index (DLQI) section than 10 and Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or and A PASI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the prior treatment course (but preferably all prior treatment courses), preferably while still on treatment but no longer following cessation of each prior treatment course The most recent PASI or DLQI assessment is no more than 1 month old at the time of application				or sole of a foot, where the plaque or plaques ere the plaques or lesions have been present ogy Life Quality Index (DLQI) score greater ced intolerable side effects from, at least three apy, methotrexate, ciclosporin, or acitretin been completed for at least the most recent e still on treatment but no longer than 1 month		
while still of face, hand severe, an	on treati d, foot, g nd for the	men jenit e fac	t but r al or f ce, pa	no longer than 1 month lexural areas at least 2 Im of a hand or sole of	of the 3 PASI symptom subscores for erythema, thic a foot the skin area affected is 30% or more of the fa an 1 month following cessation of the most recent price.	nt; for severe chronic plaque psoriasis of the ckness and scaling are rated as severe or very ce, palm of a hand or sole of a foot, as assessed

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PPLICANT (sta	amp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:	
leg No:		First Names:	First Names:	
lame:		Surname:	Surname:	
ddress:		DOB:	Address:	
		Address:		
			Fax Number:	
tanercept -	continued			
Renewal — sev	vere chronic plaque psoriasis			
Applications from	Il Number (if known): m any relevant practitioner. Approvick boxes where appropriate)			
	Following each or is sustained a or Following each	dy" severe chronic plaque psoriasis at the start of tre prior etanercept treatment course the patient has a F at this level, when compared with the pre-treatment b prior etanercept treatment course the patient has a E 5 or more, when compared with the pre-treatment ba	PASI score which is reduced by 75% or more, aseline value	
or	Following each all 3 of erythems course baseline		eduction in the PASI symptom subscores for led at this level, as compared to the treatment	
or		prior etanercept treatment course the patient has a re tained at this level, as compared to the pre-treatment		
	Patient had severe ch	ronic localised genital or flexural plaque psoriasis at t	the start of treatment	
	or Compared to the	experienced a reduction of 75% or more in the skin as pre-treatment baseline value ermatology Quality of Life Index (DLQI) improvement acing etanercept		
	·	loses no greater than 50 mg every 7 days		

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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:
Etanercept - continued		
Initial application — undifferentiated spondyloarthritis Applications only from a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)		
Patient has undifferentiated peripheral spondyloarthritis* with active peripheral joint arthritis in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip 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 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 maximum tolerated dose) and Patient 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) 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 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 Note: Indications marked with * are unapproved indications.		
Renewal — undifferentiated spondyloarthritis Current approval Number (if known):		
or Applicant is a rheumatologis Applicant is a Practitioner at continues with etanercept tr	nd confirms that a rheumatologist has provided a lette	er, email or fax recommending that the patient
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		
and Etanercept to be administered at doses no greater than 50 mg dose every 7 days		