SA2400 - Adalimumab (Amgevita)

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Ulcerative colitis - Initial application	
Ulcerative colitis - Renewal	
Undifferentiated spondyloarthritis - Initial application	
Undifferentiated spondyloarthritis - Renewal	

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APPL	ICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg N	lo:	First Names:	First Names:
Name	:	Surname:	Surname:
Addre	ss:	DOB:	Address:
		Address:	
Fax N	umber:		Fax Number:
Adal	imumab (Amgevita)		
App Prer	The patient has severe Behcet's dand The patient has severe Behcet's dand The patient has severe ocula treatment(s) appropriate for The patient has severe gastr	rointestinal, rheumatological, and/or mucocutaneous opropriate for the particular symptom(s)	not responded adequately to one or more
Note	: Indications marked with * are unapproved in	dications.	
App	al application — Hidradenitis suppurativa lications only from a dermatologist. Approvals equisites(tick boxes where appropriate)	s valid for 4 months.	
		ra Hurley Stage II or Hurley Stage III lesions in distinc	et anatomic areas
	Patient has tried, but had an inade has contraindications for systemic and	quate response to at least a 90 day trial of systemic antibiotics	antibiotics or has demonstrated intolerance to or
	Patient has 3 or more active lesion	s	
	The patient has a DLQI of 10 or m	ore and the assessment is no more than 1 month old	at time of application
Rene	ewal — Hidradenitis suppurativa		
Curre	ent approval Number (if known):		
	cations from any relevant practitioner. Approvequisites(tick boxes where appropriate)	als valid for 2 years.	
	The patient has a reduction in activand	ve lesions (e.g. inflammatory nodules, abscesses, dr	aining fistulae) of 25% or more from baseline
	The patient has a DLQI improvement	ent of 4 or more from baseline	

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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:
Adalimumab (Amgevita) - continued		
Prerequisites(tick boxes where appropriate)	vant practitioner on the recommendation of a dermate	
and	ecial Authority approval for etanercept for severe chro ed intolerable side effects	nic piaque psoriasis
or Patient has received in	nsufficient benefit to meet the renewal criteria for etan	ercept for severe chronic plaque psoriasis
or		
or present for at least 6 n	dy" severe chronic plaque psoriasis with a PASI score nonths from the time of initial diagnosis onic plaque psoriasis of the face, or palm of a hand of at least 6 months from the time of initial diagnosis	
	onic localised genital or flexural plaque psoriasis who rom the time of initial diagnosis, and with a Dermatolo	
	inadequate response to, or has experienced intolera tted doses unless contraindicated): phototherapy, me	
A PASI assessment or DLQI	assessment has been completed for at least the mo- ation of each prior treatment course and is no more th	

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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:
Renewal — Plaque psoriasis - severe chronic Current approval Number (if known):		
or The patient has experiment base or The patient has a DL or Patient had severe chronic pand The patient has experiment has experiment base or The patient has experiment has experiment base or The patient has experiment base or The patient has experiment base or The patient has experiment base or or The patient has experiment base or or The patient has experiment base or	QI improvement of 5 or more, when compared with the plaque psoriasis of the face, or palm of a hand or sole rienced reduction in the PASI symptom subscores for tained at this level, as compared to the treatment courienced reduction of 75% or more in the skin area affectaseline value	e pre-treatment baseline value e of a foot at the start of treatment all 3 of erythema, thickness and scaling, to use baseline values ected, or sustained at this level, as compared art of treatment ffected, or sustained at this level, as compared
Initial application — pyoderma gangrenosum Applications only from a dermatologist. Approvale Prerequisites(tick boxes where appropriate)	s valid without further renewal unless notified.	
	s of conventional therapy including a minimum of three d has not received an adequate response	e pharmaceuticals (e.g. prednisone, ciclosporin,

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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:	
Adalimumab (Amgevita) - continued			
Initial application — Crohn's disease - adults Applications from any relevant practitioner. Appr Prerequisites(tick boxes where appropriate)	rovals valid for 6 months.		
Patient has active Crohn's diseas	ee		
or	of greater than or equal to 300, or HBI score of greate	·	
or Patient has evidence of sho	ort gut syndrome or would be at risk of short gut sync		
Patient has an ileostomy or	r colostomy and has intestinal inflammation		
Patient has tried but had an inade and corticosteroids	equate response to, or has experienced intolerable si	de effects from, prior therapy with immunomodulators	
Renewal — Crohn's disease - adults			
Current approval Number (if known):			
Applications from any relevant practitioner. Appro			
on adalimumab	points from the CDAI score, or HBI score has reduced	d by 3 points, from when the patient was initiated	
CDAI score is 150 or less, or HBI	is 4 or less		
The patient has demonstrated an	adequate response to treatment, but CDAI score and	d/or HBI score cannot be assessed	
Initial application — Crohn's disease - childre Applications from any relevant practitioner. Appr Prerequisites(tick boxes where appropriate)			
Paediatric patient has active Crof	nn's disease		
	of greater than or equal to 30		
Patient has extensive small	I intestine disease		
Patient has tried but had an inade and corticosteroids	equate response to, or has experienced intolerable si	de effects from, prior therapy with immunomodulators	

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:				
Adalimumab (Amgevita) - continued						
Renewal — Crohn's disease - children						
Current approval Number (if known):						
Applications from any relevant practitioner. Approx						
Prerequisites(tick boxes where appropriate)						
PCDAI score has reduced by 10 po	oints from the PCDAI score when the patient was init	ated on adalimumab				
PCDAI score is 15 or less						
or	de la Constantina de					
ine patient has demonstrated an a	adequate response to treatment but PCDAI score car	inot be assessed				
Initial application — Crohn's disease - fistulisir Applications from any relevant practitioner. Appro						
Prerequisites(tick boxes where appropriate)						
Patient has confirmed Crohn's dise	ease					
Patient has one or more con	nplex externally draining enterocutaneous fistula(e)					
Patient has one or more reci	tovaginal fistula(e)					
Patient has complex peri-an	al fistula					
and A Baseline Fistula Assessment ha	and A Baseline Fistula Assessment has been completed and is no more than 1 month old at the time of application					
Renewal — Crohn's disease - fistulising						
Transmit Crommo discusso inclusing						
Current approval Number (if known):						
Applications from any relevant practitioner. Approvemental Prerequisites (tick boxes where appropriate)	<i>r</i> ais vaild for 2 years.					
The number of open draining fistul	ae have decreased from baseline by at least 50%					
	n in drainage of all fistula(e) from baseline as demon n and patient-reported pain	strated by a reduction in the Fistula Assessment				

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APPLICANT (stamp or sticker acceptable)			p or s	sticker acceptable)	PATIENT NHI:	REFERRER Reg No:	
Reg No:					First Names:	First Names:	
Name:					Surname:	Surname:	
Addre	ss: .				DOB:	Address:	
					Address:		
Fax N	umbe	ər:				Fax Number:	
Adal	imu	mab (A mg	gevita) - continued			
Appl	lication	ons from	any ı	Ocular inflammation - chrorelevant practitioner. Appropriate)			
	or	Th	ie pat	tient has had an initial Spec	cial Authority approval for infliximab for chronic ocular	inflammation	
		and	P	Patient has severe uveitis ur	ncontrolled with treatment of steroids and other immu	nosuppressants with a severe risk of vision loss	
			or [Patient is 18 years or	older and treatment with at least two other immunom	odulatory agents has proven ineffective	
			or [Patient is under 18 ye	ars and treatment with methotrexate has proven ineff	ective or is not tolerated at a therapeutic dose	
			[ars and treatment with steroids or methotrexate has p isease requires control to prevent irreversible vision l		
Rene	ewal	— Ocul	ar inf	lammation - chronic			
Curre	ent ai	oproval I	Jumb	er (if known):			
				elevant practitioner. Approv			
Prere	equis	sites(ticl	boxe	es where appropriate)			
		Tr	ie pai	tient has had a good clinica	l response following 12 weeks' initial treatment		
	or	No	omen	clature (SUN) criteria < 1/2+	period, the patient has had a sustained reduction in in anterior chamber or vitreous cells, absence of active		
	or	_ ^		macular oedema)			
					eriod, the patient has a sustained steroid sparing effe ice daily if under 18 years old	ect, allowing reduction in prednisone to < 10mg	
Appl	lication	ons from	any i	Ocular inflammation - severelevant practitioner. Appro			
	or	☐ Pa	atient	has had an initial Special A	Authority approval for infliximab for severe ocular infla	mmation	
		and] P	atient has severe, vision-th	reatening ocular inflammation requiring rapid control		
			or [Treatment with high-d ineffective at controllir	ose steroids (intravenous methylprednisolone) followe	ed by high dose oral steroids has proven	
			[Patient developed nev	v inflammatory symptoms while receiving high dose s	steroids	
			or [Patient is aged under ineffective at controllir	8 years and treatment with high dose oral steroids and symptoms	nd other immunosuppressants has proven	

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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:		
Fax Num	ber:						Fax Number:
Adalim	umab ((Am	ngevit	ta) - continued			
Renewa	I — Ocu	lar i	nflamm	nation - severe			
Current	approval	Nun	nber (if	known):			
			•	unt practitioner. Appro			
Prerequ	i isites (tio	ck bo	xes wh	nere appropriate)			
		he p	atient h	has had a good clinica	al response following 3 initial do	oses	
or		ollov	wing ea	ach 2 year treatment	period, the patient has had a su	ustained reduction in in	nflammation (Standardisation of Uveitis
				re (SUN) criteria < ½+ ular oedema)	anterior chamber or vitreous of	cells, absence of active	vitreous or retinal lesions, or resolution of uveitic
or		ollov	wing ea	ach 2 year treatment p	eriod, the patient has a sustair	ned steroid sparing effe	ect, allowing reduction in prednisone to < 10mg
					ice daily if under 18 years old		, ,
Initial a	oplicatio	n —	ankvlo	osing spondylitis			
Applicat	tions only	y froi	m a rhe	eumatologist. Approva nere appropriate)	als valid for 6 months.		
Ticicqu			7,00 WII				
	and		Patien	nt has had an initial Sp	ecial Authority approval for eta	nercept for ankylosing	spondylitis
	and			The natient has eyne	rienced intolerable side effects		
		or				Ale a vere event evite vie few	
				The patient has received	ved insufficient benefit to meet	the renewal chierla for	ankylosing spondylitis
or		_					
	and		Patien	it has a confirmed dia	gnosis of ankylosing spondylitis	s for more than six mor	nths
	and		Patien	nt has low back pain a	nd stiffness that is relieved by e	exercise but not by rest	t
	and		Patien	nt has bilateral sacroili	tis demonstrated by radiology i	imaging	
	anu					o or more NSAIDs, whi	ile patient was undergoing at least 3 months of
	and		a regu	ılar exercise regimen f	or ankylosing spondylitis		
							rontal planes as determined by the following and lumbar side flexion measurement of less
		or		than or equal to 10 cr	n (mean of left and right)		
				Patient has limitation gender	of chest expansion by at least 2	2.5 cm below the avera	age normal values corrected for age and
	and	<u> </u>					
		Ш			0-10 scale completed after the an 1 month old at the time of a		but prior to ceasing any previous pharmacological
1							

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APPLICANT (stamp or sticker acceptable)			PATIENT NHI:	REFERRER Reg No:	
Reg No:			First Names:	First Names:	
Name:			Surname:	Surname:	
Addre	ess:		DOB:	Address:	
			Address:		
Fax N	lumbei	,. 		Fax Number:	
Adal	imun	nab (Amgevita) - continued			
Ren	ewal –	- ankylosing spondylitis			
Curr	ent ap	proval Number (if known):			
Appl	ication	s from any relevant practitioner. Appro			
Prer	equisi —	tes(tick box where appropriate)			
Į		reatment has resulted in an improveme ASDAI of 50%, whichever is less	ent in BASDAI of 4 or more points from pre-treatment	baseline on a 10 point scale, or an improvement in	
Initis	al anni	ication — Arthritis - oligoarticular c	ourse iuvenile idionathic		
App	lication	ns only from a named specialist or rheu	umatologist. Approvals valid for 6 months.		
Prer	equisi	tes(tick boxes where appropriate)			
		The patient has had an initi	al Special Authority approval for etanercept for oligoa	rticular course juvenile idiopathic arthritis (JIA)	
		and			
	or Patient has experience		ced intolerable side effects		
		Patient has received	insufficient benefit to meet the renewal criteria for olig	poarticular course JIA	
	or				
		To be used as an adjunct to	methotrexate therapy or monotherapy where use of	methotrexate is limited by toxicity or intolerance	
			ar course JIA for 6 months duration or longer		
		and			
		maximum tolerated d	s with limited range of motion, pain or tenderness aftence)	er a 3-month trial of methotrexate (at the	
			ease activity (cJADAS10 score greater than 1.5) with	poor prognostic features after a 3-month trial of	
		methotrexate (at the i	maximum tolerated dose)		
Rene	ewal –	- Arthritis - oligoarticular course juv	venile idiopathic		
Curre	ent ap _l	proval Number (if known):			
		s from any relevant practitioner. Appro	vals valid for 2 years.		
Prer	equisi	tes(tick boxes where appropriate)			
	or [Following initial treatment, the pat assessment from baseline	ient has at least a 50% decrease in active joint count	and an improvement in physician's global	
] <u> </u>	On subsequent reapplications, the improvement in physician's global	e patient demonstrates at least a continuing 30% impassessment from baseline	rovement in active joint count and continued	

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)			le) PATIENT NHI:	REFERRER Reg No:
Reg No:			First Names:	First Names:
Name:	:		Surname:	Surname:
Addres	ss:		DOB:	Address:
			Address:	
Fax N	umber	······································		Fax Number:
Adali	mun	nab (Amgevita) - contin	nued	
Appli	ication	Patient has had a and Patient has had a and Patient has had a Patient has Or Patient has At least 5 a methotrexa Or Moderate of tolerated do	an initial Special Authority approval for etanercep sexperienced intolerable side effects areceived insufficient benefit to meet the renewal adjunct to methotrexate therapy or monotherapy polyarticular course JIA for 6 months duration or lactive joints and at least 3 joints with limited rangulate (at the maximum tolerated dose) or high disease activity (cJADAS10 score of at least	t for polyarticular course juvenile idiopathic arthritis (JIA) I criteria for polyarticular course JIA where use of methotrexate is limited by toxicity or intolerance longer ge of motion, pain or tenderness after a 3-month trial of ast 2.5) after a 3-month trial of methotrexate (at the maximum
Rene	wal –	– Arthritis - polyarticular c	ourse juvenile idiopathic	
Applic	cations	s from any relevant practition tes(tick boxes where approp Following initial treatme assessment from baseli On subsequent reapplic	ent, the patient has at least a 50% decrease in ac ine	ctive joint count and an improvement in physician's global nuing 30% improvement in active joint count and continued

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)			mp o	r sticker acceptable)	PATIENT NHI:	REFERRER Reg No:	
Reg No:					. First Names:	First Names:	
Name:					. Surname:	Surname:	
Addre	ess:				. DOB:	Address:	
					. Address:		
						Fax Number:	
Initia App	al app	lication	on — ly froi	Arthritis - psoriatic m a rheumatologist. Appropriate where appropriate)	vals valid for 6 months.		
		and		Patient has had an initial	Special Authority approval for etanercept or secukinum	nab for psoriatic arthritis	
			or	The patient has exp	erienced intolerable side effects		
				The patient has rec	eived insufficient benefit from to meet the renewal crite	eria for psoriatic arthritis	
or		and and and	Patient has tried and not responded to at least three months Patient has tried and not responded to at least three months contraindicated) and Patient has persistent symptoms of poorly controlled a elbow, knee, ankle, and either shoulder or hip and		esponded to at least three months of methotrexate at esponded to at least three months of sulfasalazine or nt symptoms of poorly controlled and active disease in the symptoms of poorly controlled and active disease in	ns of methotrexate at a maximum tolerated dose (unless contraindicated) ns of sulfasalazine or leflunomide at maximum tolerated doses (unless and active disease in at least 15 swollen joints and active disease in at least four joints from the following: wrist,	
			or		greater than 25 mm per hour		
					neasured as patient is currently receiving prednisone more than three months	therapy at a dose of greater than 5 mg per day	
Rene	ewal –	– Art	hritis	- psoriatic			
Appli	ication	s fror	n any	nber (if known): r relevant practitioner. App oxes where appropriate)	rovals valid for 2 years.		
	0,			wing initial treatment, the ponse in the opinion of the p	atient has at least a 50% decrease in swollen joint counysician	unt from baseline and a clinically significant	
	or 			nt demonstrates at least a pinion of the treating physic	continuing 30% improvement in swollen joint count fro cian	m baseline and a clinically significant response in	

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APPLICANT (stamp or sticker acceptable)			o or	sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:					First Names:	First Names:
Name:					Surname:	Surname:
Addre	ss:				DOB:	Address:
					Address:	
Fax N	umbei	r:				Fax Number:
Adali	imun	nab (A	m	gevita) - continued		
Appl	lication	ns only f	rom	Arthritis - rheumatoid a rheumatologist. Approva ses where appropriate)	lls valid for 6 months.	
		and] ·	The patient has had an initia	al Special Authority approval for etanercept for rheum	atoid arthritis
			or		rienced intolerable side effects ved insufficient benefit from etanercept to meet the re	newal criteria for rheumatoid arthritis
	or					
or		Patient has had rheumatoid arthritis (either confirmed by radiology immonths duration or longer and Treatment is to be used as an adjunct to methotrexate therapy or monintolerance and Patient has tried and not responded to at least three months of methore sulphate at maximum tolerated doses (unless contraindicated) Patient has tried and not responded to at least three months of dose of ciclosporin Patient has tried and not responded to at least three months of dose of ciclosporin Patient has tried and not responded to at least three months of alone or in combination with methotrexate and Patient has persistent symptoms of poorly controlled and active elbow, knee, ankle, and either shoulder or hip			an adjunct to methotrexate therapy or monotherapy when adjunct to methotrexate therapy or monotherapy when a ponded to at least three months of methotrexate in content doses (unless contraindicated) not responded to at least three months of methotrexate and responded to at least three months of therapy at the polymethod with methotrexate and active disease in symptoms of poorly controlled and active disease in	here use of methotrexate is limited by toxicity or maximum tolerated dose (unless contraindicated) ombination with sulfasalazine and hydroxychloroquir te in combination with the maximum tolerated the maximum tolerated dose of leflunomide at least 15 swollen joints
Rene	ewal –	– Arthri	tis	- rheumatoid		
Appli	cation	s from a	ıny	ber (if known): relevant practitioner. Approves where appropriate)		
	or [ing initial treatment, the pati	ent has at least a 50% decrease in active joint count on of the physician	from baseline and a clinically significant
	[patient demonstrates at least a continuing 30% impresatment in the opinion of the physician	ovement in active joint count from baseline and a

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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:					
Adal	Adalimumab (Amgevita) - continued								
App	Initial application — Still's disease - adult-onset (AOSD) Applications only from a rheumatologist. Approvals valid without further renewal unless notified. Prerequisites(tick boxes where appropriate)								
		The patient has had an initial Special Authority approval for etanercept and/or tocilizumab for AOSD Patient has experienced intolerable side effects from etanercept and/or tocilizumab or							
	or	Patient has received insufficient benefit from at least a three-month trial of etanercept and/or tocilizumab							
		Patient diagnosed with AOSD according to the Yamaguchi criteria Patient has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, NSAIDs and methotrexate Patient has persistent symptoms of disabling poorly controlled and active disease							
Initial application — ulcerative colitis Applications from any relevant practitioner. Approvals valid for 3 months. Prerequisites(tick boxes where appropriate)									
	and	Patient has active ulcerative colitis							
		Patient's SCCAI score is gre	eater than or equal to 4						
		Patient's PUCAI score is greater than or equal to 20							
	and	Patient has tried but had an inadequate response to, or has experienced intolerable side effects from prior therapy with immunomodulators and systemic corticosteroids Surgery (or further surgery) is considered to be clinically inappropriate							
		Surgery (or further surgery) is con	sidered to be clinically mappropriate						
Rene	ewal –	– ulcerative colitis							
Appli	Current approval Number (if known):								
	ا	The SCCAI score has reduced by	by 2 points or more from the SCCAI score when the patient was initiated on biologic therapy						
	or 	The PUCAI score has reduced by	10 points or more from the PUCAI score when the pa	tient was initiation on biologic therapy					

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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:						
Addres	SS:	DOB:	Address:						
		Address:							
Fax N	umber:		Fax Number:						
Adali	Adalimumab (Amgevita) - continued								
Initial application — undifferentiated spondyloarthritis Applications only from a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)									
	Patient has undifferentiated periph wrist, elbow, knee, ankle, and either and	eral spondyloarthritis* with active peripheral joint arther shoulder or hip	nritis in at least four joints from the following:						
	Patient has tried and not responded to at least three months of each of methotrexate, sulfasalazine and leflunomide, at maxim tolerated doses (unless contraindicated) and								
	Patient has a CRP level great	ater than 15 mg/L measured no more than one month prior to the date of this application							
		Patient has an ESR greater than 25 mm per hour measured no more than one month prior to the date of this application							
		d as patient is currently receiving prednisone therapy months	at a dose of greater than 5 mg per day and has						
Note:	Indications marked with * are unapproved in	dications							
Rene	wal — undifferentiated spondyloarthritis								
C	at approval Number (if Iraqua)								
	nt approval Number (if known): cations from any relevant practitioner. Approv								
	equisites(tick boxes where appropriate)	•							
	response to treatment in the opinion	ent has at least a 50% decrease in active joint count on of the physician	from baseline and a clinically significant						
	The patient demonstrates at least in the opinion of the treating physic	a continuing 30% improvement in active joint count fr cian	om baseline and a clinically significant response						
Initial application — inflammatory bowel arthritis – axial Applications only from a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)									
	Patient has a diagnosis of active u	Icerative colitis or active Crohn's disease							
	Patient has axial inflammatory pair	n for six months or more							
	Patient is unable to take NSAIDs								
		demonstrated by radiological imaging or MRI							
	physiotherapist	tely to prior treatment consisting of at least 3 months	s of an exercise regime supervised by a						
	A BASDAI of at least 6 on a 0-10 treatment	scale completed after the 3 month exercise trial, but	prior to ceasing any previous pharmacological						

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APPLICANT (stamp or sticker acceptable)		PATIENT NHI:	REFERRER Reg No:
Reg N	No:	First Names:	First Names:
Name	2:	Surname:	Surname:
Address:		DOB:	Address:
		Address:	
Fax N	lumber:		Fax Number:
Adal	imumab (Amgevita) - continued		
Rene	ewal — inflammatory bowel arthritis – axia	I	
Curr	ent approval Number (if known):		
Appl	ications from any relevant practitioner. Appro		
Prer	equisites(tick box where appropriate)		
[Treatment has resulted in an improveme BASDAI of 50%, whichever is less	nt in BASDAI of 4 or more points from pre-treatment	baseline on a 10 point scale, or an improvement in
App Prem	and Patient has active arthritis in at leasternoclavicular Patient has tried and not experience (unless contraindicated) and Patient has tried and not experience contraindicated) and Patient has tried and not experience contraindicated) and Patient has a CRP level greence or Patient has an ESR greater or ESR and CRP not measure done so for more than three	elcerative colitis or active Crohn's disease ast four joints from the following: hip, knee, ankle, subset of a response to at least three months of methotrex aced a response to at least three months of sulfasalazed ater than 15 mg/L measured no more than one month than 25 mm per hour measured no more than one mode as patient is currently receiving prednisone therapy months	ate, or azathioprine at a maximum tolerated dose ine at a maximum tolerated dose (unless in prior to the date of this application onth prior to the date of this application
Curro Appl	ewal — inflammatory bowel arthritis – peri ent approval Number (if known): ications from any relevant practitioner. Appro equisites(tick boxes where appropriate)	· ·····	
	treatment in the opinion of the phy	has at least a 50% decrease in active joint count fror sician	n baseline and a clinically significant response to
	Patient has experienced at least a physician	continuing 30% improvement in active joint count fro	m baseline in the opinion of the treating