SA2179 - Infliximab

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Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab		
or Patient has extensive small i or Patient has evidence of shor or Patient has an ileostomy or or		ome with further bowel resection
or initiated on infliximab CDAI score is 150 or less, or The patient has demonstrate and Infliximab to be administered at do to 3 doses if required for secondar	rals valid for 2 years. 100 points from the CDAI score, or HBI score has red	e and/or HBI score cannot be assessed very 8 weeks (or equivalent) can be used for up er re-induction may be considered sixteen weeks

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Address:	DOB:	Address:	
	Address:		
Fax Number:		Fax Number:	
Infliximab - continued			
Initial application — Crohn's disease (children Applications from any relevant practitioner. Appro Prerequisites(tick boxes where appropriate)			
Paediatric patient has active Croh	n's disease		
	of greater than or equal to 30		
or Patient has extensive small	intestine disease		
and Patient has extensive small mestine disease and Patient has tried but experienced an inadequate response to, or intolerable side effects from, prior therapy with immunomodulators and corticosteroids			
Current approval Number (if known):			
or PCDAI score has reduced b			
or The patient has demonstrate			
Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle. Up to 10 mg/kg every 8 weeks (or equivalent) may be used for patients treated with this dose prior to 1 February 2019			
Initial application — Graft vs host disease Applications from any relevant practitioner. Appro Prerequisites(tick box where appropriate)	vals valid without further renewal unless notified.		
Patient has steroid-refractory acute graft	vs. host disease of the gut		
Initial application — Pulmonary sarcoidosis Applications from any relevant practitioner. Appro Prerequisites(tick box where appropriate)	evals valid without further renewal unless notified.	is refractory to other treatments	

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Name:	Surname:	Surname:	
Address:	DOB:	Address:	
	Address:		
Fax Number:		Fax Number:	
Infliximab - continued			
Initial application — acute fulminant ulcerative colitis Applications only from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 6 weeks. Prerequisites(tick boxes where appropriate) Patient has acute, fulminant ulcerative colitis and Treatment with intravenous or high dose oral corticosteroids has not been successful			
Initial application — ankylosing spondylitis Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 3 months. Prerequisites(tick boxes where appropriate) The patient has had an initial Special Authority approval for adalimumab and/or etanercept for ankylosing spondylitis The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept or Following 12 weeks of adalimumab and/or etanercept treatment, the patient did not meet the renewal criteria for adalimumab and/or etanercept for ankylosing spondylitis			
Renewal — ankylosing spondylitis			
Current approval Number (if known):			

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Reg No:	First Names:	First Names:
Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
or The patient has exper Or Patient has severe uveitis ur Or Patient is 18 years or Or Patient is under 18 ye Or Patient is under 8 years		enewal criteria for adalimumab for chronic nosuppressants with a severe risk of vision loss odulatory agents has proven ineffective ective or is not tolerated at a therapeutic dose roven ineffective or is not tolerated at a
Renewal — chronic ocular inflammation Current approval Number (if known):	vals valid for 12 months.	
The patient has had a good clinical response following 3 initial doses or Following each 12 month treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < ½+ anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uvei cystoid macular oedema) or Following each 12 month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10r		vitreous or retinal lesions, or resolution of uveitic
Note: A trial withdrawal should be considered afte vision loss if infliximab is withdrawn.	r every 24 months of stability, unless the patient is de	emed to have extremely high risk of irreversible

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Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
Initial application — fistulising Crohn's disease Applications from any relevant practitioner. Appro Prerequisites (tick boxes where appropriate) Patient has confirmed Crohn's disease (appropriate) Patient has one or more confirmed or Patient has one or more reconfirmed Crohn's disease (appropriate)	ease nplex externally draining enterocutaneous fistula(e) tovaginal fistula(e)	
Prerequisites(tick boxes where appropriate) The number of open draining or There has been a marked re	g fistulae have decreased from baseline by at least 5 eduction in drainage of all fistula(e) from baseline (in the sessment score), together with less induration and pati	0% he case of adult patients, as demonstrated by a
Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle. Up to 10 mg/kg every 8 weeks (or equivalent) may be used for patients treated with this dose prior to 1 February 2019		
Initial application — neurosarcoidosis Applications only from a neurologist or Practitione Prerequisites(tick boxes where appropriate)	er on the recommendation of a neurologist. Approvals	s valid for 18 months.
and Patient has CNS involvement and Patient has steroid-refractory disea		
IV cyclophosphamide has been tried or		
Treatment with IV cyclophos	phamide is clinically inappropriate	

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Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
Current approval Number (if known): Applications only from a neurologist or Practitioner Prerequisites(tick boxes where appropriate)	on the recommendation of a neurologist. Approvals	valid for 18 months.
A withdrawal period has been tried	and the patient has relapsed	
A withdrawal period has been considered but would not be clinically appropriate		
There has been a marked re	duction in prednisone dose	
There has been an im	provement in MRI appearances	
Marked improvement i	n other symptomology	

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APPLICANT (stamp or s	sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:		First Names:	First Names:
Name:		Surname:	Surname:
Address:		DOB:	Address:
		Address:	
Fax Number:			Fax Number:
Infliximab - continued	d		
Prerequisites(tick boxe	a dermatologist or Practitices where appropriate) the patient has had an initial soriasis Patient has experience Patient has received in adalimumab, etanerose Patient has "whole book greater than 10, where the patient has severe check the patie	al Special Authority approval for adalimumab, etanerous intolerable side effects from adalimumab, etanerous or secutions and alimumab, etanerous or secutions and alimumab, etanerous or secution adalimumab, etanerous or secution adalimumab or secuti	ept or secukinumab for severe chronic plaque ept or secukinumab ecukinumab to meet the renewal criteria for is Area and Severity Index (PASI) score of m the time of initial diagnosis
and A and T Note: "Inadequate resp while still on treatment thand or foot, at least 2	f the following (at maximun PASI assessment has been ourses), preferably while state the most recent PASI assessment in the most recent in the most recent path in th	inadequate response (see Note) to, or has experience to tolerated doses unless contraindicated): phototheral en completed for at least the most recent prior treatmential on treatment but no longer than 1 month following essment is no more than 1 month old at the time of initial tole body severe chronic plaque psoriasis, a PASI sort following cessation of the most recent prior treatment becomes for erythema, thickness and scaling are rated dor sole of a foot, as assessed preferably while still of	apy, methotrexate, ciclosporin, or acitretin ent course (but preferably all prior treatment cessation of each prior treatment course liation ore of greater than 10, as assessed preferably at; for severe chronic plaque psoriasis of the face, d as severe or very severe, and the skin area

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Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
Renewal — plaque psoriasis		
Prerequisites(tick boxes where appropriate) Patient had "whole box and Following each prior in	dy" severe chronic plaque psoriasis at the start of treatment course the patient has a PASI scowhen compared with the pre-infliximab treatment bases	atment re which is reduced by 75% or more, or is
Patient had severe chr	ronic plaque psoriasis of the face, or palm of a hand of	or sole of a foot at the start of treatment
all 3 of erythems course baseline or Following each p	prior infliximab treatment course the patient has a rec a, thickness and scaling, to slight or better, or sustaine values prior infliximab treatment course the patient has a rec ained at this level, as compared to the pre-infliximab	ed at this level, as compared to the treatment luction of 75% or more in the skin area
and Infliximab to be administered at dos	ses no greater than 5 mg/kg every 8 weeks	

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Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
Initial application — previous use Applications from any relevant practitioner. Appro Prerequisites(tick boxes where appropriate) Patient was being treated with infliand Rheumatoid arthritis Or Ankylosing spondylitis Or Severe ocular inflammation Or Chronic ocular inflammation Or Crohn's disease (adults) Or Fistulising Crohn's disease Or Severe fulminant ulcerative Or Severe ulcerative colitis Or Plaque psoriasis Or Neurosarcoidosis Or Severe Behcet's disease	ximab prior to 1 February 2019	
Initial application — psoriatic arthritis Applications only from a rheumatologist or Practiti Prerequisites(tick boxes where appropriate)	ioner on the recommendation of a rheumatologist. Ap	oprovals valid for 4 months.
The patient has had an initial Spec	cial Authority approval for adalimumab and/or etanero	ept and/or secukinumab for psoriatic arthritis
The patient has experienced or	d intolerable side effects from adalimumab and/or eta	nercept and/or secukinumab
Following 3-4 months' initial	treatment with adalimumab and/or etanercept and/or nab and/or etanercept and/or secukinumab for psoria	

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Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Infliximab - continued		
Renewal — psoriatic arthritis		
Current approval Number (if known):		
• • • • • • • • • • • • • • • • • • • •	oner on the recommendation of a rheumatologist. Ap	provals valid for 6 months.
Prerequisites(tick boxes where appropriate)		
	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
The patient demonstrates a	least a continuing 30% improvement in active joint or treatment in the opinion of the treating physician	ount from baseline and a clinically significant
and Infliximab to be administered at do	ses no greater than 5 mg/kg every 8 weeks	
Initial conditions who we staid authorities		
Initial application — rheumatoid arthritis Applications only from a rheumatologist or Practit Prerequisites(tick boxes where appropriate)	ioner on the recommendation of a rheumatologist. Ap	oprovals valid for 4 months.
The patient has had an initial Spec	cial Authority approval for adalimumab and/or etanero	ept for rheumatoid arthritis
The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept		
Following at least a four month trial of adalimumab and/or etanercept, the patient did not meet the renewal criteria for adalimumab and/or etanercept		
and Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance		
Renewal — rheumatoid arthritis		
0		
Current approval Number (if known):	oner on the recommendation of a rheumatologist. Ap	provals valid for 6 months.
Prerequisites(tick boxes where appropriate)	ğ ,	•
Treatment is to be used as an adjuintolerance	unct to methotrexate therapy or monotherapy where u	se of methotrexate is limited by toxicity or
Following 3 to 4 months' in clinically significant respons	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
The patient demonstrates a response to treatment in the	least a continuing 30% improvement in active joint coopinion of the physician	ount from baseline and a clinically significant
and Infliximab to be administered at doses no greater than 3 mg/kg every 8 weeks		

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Name:	Surname:	Surname:	
Address:	DOB:	Address:	
	Address:		
Fax Number:		Fax Number:	
Infliximab - continued			
Initial application — severe Behcet's disease Applications from any relevant practitioner. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate) The patient has severe Behcet's disease which is significantly impacting the patient's quality of life (see Notes) The patient has severe ocular, neurological and/or vasculitic symptoms and has not responded adequately to one or more treatment(s) appropriate for the particular symptom(s) (see Notes) The patient has severe gastrointestinal, rheumatologic and/or mucocutaneous symptoms and has not responded adequately to two or more treatment appropriate for the particular symptom(s) (see Notes) and The patient is experiencing significant loss of quality of life Note: Behcet's disease diagnosed according to the International Study Group for Behcet's Disease. Lancet 1990;335(8697):1078-80. Quality of life measured using an appropriate quality of life scale such as that published in Gilworth et al J Rheumatol. 2004;31:931-7. Treatments appropriate for the particular symptoms are those that are considered standard conventional treatments for these symptoms, for example intravenous/oral steroids and other immunosuppressants for ocular symptoms; are those that are considered standard conventional treatments for these symptoms, for example intravenous/oral steroids and other immunosuppressants for ocular symptoms; seroids, thalidomide, interferon alpha and ciclosporin for mucocutaneous symptoms; and colchicine, steroids and methotrexate for rheumatological symptoms.			
Renewal — severe Behcet's disease			
Current approval Number (if known):			
Applications from any relevant practitioner. Approx	vals valid for 6 months.		
Prerequisites(tick boxes where appropriate)			
Patient has had a good clinical res	sponse to initial treatment with measurably improved of	quality of life	
	Infliximab to be administered at doses no greater than 5 mg/kg every 8 weeks		
Renewal — fulminant ulcerative colitis			
Current approval Number (if known):			
Applications from any relevant practitioner. Approx Prerequisites (tick boxes where appropriate)			
Where maintenance treatment is or reassessed every 6 months	considered appropriate, infliximab should be used in c	combination with immunomodulators and	
Infliximab to be administered at do to 3 doses if required for secondar	uses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every non-response to treatment for re-induction. Another on cycle. Up to 10 mg/kg every 8 weeks (or equivaler	r re-induction may be considered sixteen weeks	

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Name:	Surname:	Surname:		
Address:	DOB:	Address:		
	Address:			
Fax Number:		Fax Number:		
Infliximab - continued				
or The patient has experience or The patient has receive ocular inflammation Or Patient has severe, vision-the ineffective at controllin or Patient developed new or	vals valid for 4 months. I Special Authority approval for adalimumab for sever enced intolerable side effects from adalimumab ed insufficient benefit from adalimumab to meet the reatening ocular inflammation requiring rapid control ose steroids (intravenous methylprednisolone) followed symptoms v inflammatory symptoms while receiving high dose self-search and treatment with high dose oral steroids are	enewal criteria for adalimumab for severe ed by high dose oral steroids has proven teroids		
Renewal — severe ocular inflammation Current approval Number (if known):				
Nomenclature (SUN) criteria < ½+ cystoid macular oedema) or Following each 12 month treatmen	nt period, the patient has had a sustained reduction in anterior chamber or vitreous cells, absence of active t period, the patient has a sustained steroid sparing e	vitreous or retinal lesions, or resolution of uveitic		
daily, or steroid drops less than twice daily if under 18 years old Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely high risk of irreversible vision loss if infliximab is withdrawn.				

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Name:	Surname:	Surname:		
Address:	DOB:	Address:		
	Address:			
Fax Number:		Fax Number:		
Initial application — ulcerative colitis Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate) Patient has active ulcerative colitis and Patients SCCAI is greater than or equal to 4 or Patients PUCAI score is greater than or equal to 20 and Patient has tried but has experienced an inadequate response to, or has experienced intolerable side effects from, prior therapy with immunomodulators and systemic corticosteroids				
Current approval Number (if known):				
The SCCAI score has reduced by 2 points or more from the SCCAI score when the patient was initiated on infliximab The PUCAI score has reduced by 10 points or more from the PUCAI score when the patient was initiated on infliximab Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle. Up to 10 mg/kg every 8 weeks (or equivalent) may be used for patients treated with this dose prior to 1 February 2019				
Initial application — pyoderma gangrenosum Applications only from a dermatologist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)				
	of conventional therapy including a minimum of three not received an adequate response	e pharmaceuticals (e.g. prednisone, ciclosporine,		
Note: Note: Indications marked with * are unappro	oved indications.			

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Address:	DOB:	Address:			
	Address:				
Fax Number:		Fax Number:			
Infliximab - continued					
Renewal — pyoderma gangrenosum					
Current approval Number (if known):					
	ner on the recommendation of a dermatologist. Appro	ovals valid for 4 months.			
Prerequisites(tick boxes where appropriate)					
Patient has shown clinical improvement					
and Patient continues to require treatm	ent				
and A maximum of 8 doses					
A maximum of 6 doses					
Initial application — inflammatory bowel arthrit					
Applications from any relevant practitioner. Appro Prerequisites (tick boxes where appropriate)	vals valid for 6 months.				
D. Bullium have a discount of a city of	Contract of the contract of Contracts of the contract				
and	Icerative colitis or active Crohn's disease				
Patient has had axial inflammatory	pain for six months or more				
Patient is unable to take NSAIDs					
	Patient has unequivocal sacroiliitis demonstrated by radiological imaging or MRI and Patient's disease has not responded adequately to prior treatment consisting of at least 3 months of an exercise regime supervised by a physiotherapist				
Patient's disease has not responde physiotherapist					
Patient has a BASDAI of at least 6 pharmacological treatment	on a 0 - 10 scale completed after the 3 month exer	cise trial, but prior to ceasing any previous			
Renewal — inflammatory bowel arthritis – axial					
Current approval Number (if known):					
Applications from any relevant practitioner. Approvals valid for 2 years.					
Prerequisites(tick box where appropriate)					
Treatment has resulted in an improvement in BASDAI of 4 or more points from pre-treatment baseline on a 10-point scale, or an improvement in BASDAI of 50%, whichever is less					

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Name:		Surname:	Surname:	
Addres	s:	DOB:	Address:	
		Address:		
Fax Nu	mber:		Fax Number:	
Inflixi	mab - continued			
Initial application — inflammatory bowel arthritis – peripheral Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate) Patient has a diagnosis of active ulcerative colitis or active Crohn's disease and Patient has active arthritis in at least four joints from the following: hip, knee, ankle, subtalar, tarsus, forefoot, wrist, elbow, shoulder, sternoclavicular Patient has tried and not experienced a response to at least three months of methotrexate or azathioprine at a maximum tolerated dose (unless contraindicated) Patient has tried and not experienced a response to at least three months of sulfasalazine at a maximum tolerated dose (unless contraindicated) Patient has a CRP level greater 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 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				
Renewal — inflammatory bowel arthritis – peripheral				
Current approval Number (if known):				
Applications from any relevant practitioner. Approvals valid for 2 years. Prerequisites(tick boxes where appropriate)				
	or response to treatment in the opinion	has experienced at least a 50% decrease in active jo on of the physician continuing 30% improvement in active joint count fro		