## SA2402 - Infliximab

Crohn's disease (adults) - Initial application
Crohn's disease (adults) - Renewal
Crohn's disease (children) - Initial application
Crohn's disease (children) - Renewal
Graft vs host disease - Initial application
Pulmonary sarcoidosis - Initial application
Acute fulminant ulcerative colitis - Initial application
Ankylosing spondylitis - Initial application
Ankylosing spondylitis - Renewal
Chronic ocular inflammation - Initial application
Chronic ocular inflammation - Renewal
Fistulising Crohn's disease - Initial application
Fistulising Crohn's disease - Renewal
Fulminant ulcerative colitis - Renewal
Inflammatory bowel arthritis – axial - Initial application
Inflammatory bowel arthritis – axial - Renewal
Inflammatory bowel arthritis – peripheral - Initial application
Inflammatory bowel arthritis - peripheral - Renewal
Neurosarcoidosis - Initial application
Neurosarcoidosis - Renewal
Plaque psoriasis - Initial application
Plaque psoriasis - Renewal
Previous use - Initial application
Psoriatic arthritis - Initial application
Psoriatic arthritis - Renewal
Pyoderma gangrenosum - Initial application
Pyoderma gangrenosum - Renewal
Rheumatoid arthritis - Initial application11
Rheumatoid arthritis - Renewal
Severe Behcet's disease - Initial application
Severe Behcet's disease - Renewal
Severe ocular inflammation - Initial application
Severe ocular inflammation - Renewal
Ulcerative colitis - Initial application
Ulcerative colitis - Renewal

APPLICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:	First Names:	First Names:
Name:	Surname:	Surname:
	DOB:	
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Fax Number:		Fax Number:

## Infliximab

Appli	catior	<b>cation — Crohn's disease (adults)</b> s from any relevant practitioner. Approvals valid for 6 months. <b>es</b> (tick boxes where appropriate)	
	and	Patient has active Crohn's disease	
		Patient has a CDAI score of greater than or equal to 300 or HBI score of greater than or equal to 10 or	
		Patient has extensive small intestine disease affecting more than 50 cm of the small intestine <b>or</b>	
		Patient has evidence of short gut syndrome or would be at risk of short gut syndrome with further bowel resection or	
		Patient has an ileostomy or colostomy, and has intestinal inflammation	
	and [	Patient has tried but has experienced an inadequate response to, or intolerable side effects from, prior therapy with immunomodulators and corticosteroids	
Rene	wal –	Crohn's disease (adults)	
Appli	cation	roval Number (if known): from any relevant practitioner. Approvals valid for 2 years. <b>es</b> (tick boxes where appropriate)	
		or CDAI score has reduced by 100 points from the CDAI score, or HBI score has reduced by 3 points, from when the patient was initiated on infliximab CDAI score is 150 or less, or HBI is 4 or less	
		or The patient has demonstrated an adequate response to treatment but CDAI score and/or HBI score cannot be assessed	
	and [	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	

Enquiries	to Ministry	of Health
0800 855	066	

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Infliximab - continued

Initial application — Crohn's disease (children) Applications from any relevant practitioner. Approvals valid for 6 months.	
Prerequisites(tick boxes where appropriate)	
Paediatric patient has active Crohn's disease	
Patient has a PCDAI score of greater than or equal to 30	
Patient has extensive small intestine disease	
and Patient has tried but experienced an inadequate response to, or intolerable side effects from, prior therapy with immunomodulators are corticosteroids	nd
Renewal — Crohn's disease (children)	
Current approval Number (if known):	
Applications from any relevant practitioner. Approvals valid for 2 years.	
Prerequisites (tick boxes where appropriate)	
PCDAI score has reduced by 10 points from the PCDAI score when the patient was initiated on infliximab	
PCDAI score is 15 or less	
or The patient has demonstrated an adequate response to treatment but PCDAI score cannot be assessed	
and 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 u to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen wee 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 d prior to 1 February 2019	ks
Initial application — Graft vs host disease Applications from any relevant practitioner. Approvals valid without further renewal unless notified. Prerequisites(tick box where appropriate)	
Patient has steroid-refractory acute graft vs. host disease of the gut	
Initial application — Pulmonary sarcoidosis Applications from any relevant practitioner. Approvals valid without further renewal unless notified. Prerequisites(tick box where appropriate)	
Patient has life-threatening pulmonary sarcoidosis diagnosed by a multidisciplinary team that is refractory to other treatments	

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	First Names:

### 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
and 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
Ponowal ankyloging spondylitis
Renewal — ankylosing spondylitis

Current approval Number (if known):.....

Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)

> Following 12 weeks of infliximab treatment, BASDAI has improved by 4 or more points from pre-infliximab baseline on a 10 point scale, or by 50%, whichever is less

Physician considers that the patient has benefited from treatment and that continued treatment is appropriate

Infliximab to be administered at doses no greater than 5 mg/kg every 6-8 weeks

I confirm the above details are correct and that in signing this form I understand I may be audited.

and

and

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### Infliximab - continued

oplicatio	ns fron	n — chronic ocular inflammation n any relevant practitioner. Approvals valid for 4 months. k boxes where appropriate)
	and	The patient has had an initial Special Authority approval for adalimumab for chronic ocular inflammation
		The patient has experienced intolerable side effects from adalimumab
		or The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for chronic ocular inflammation
or	and	Patient has severe uveitis uncontrolled with treatment of steroids and other immunosuppressants with a severe risk of vision loss
		Patient is 18 years or older and treatment with at least two other immunomodulatory agents has proven ineffective or
		Patient is under 18 years and treatment with methotrexate has proven ineffective or is not tolerated at a therapeutic dose or
		Patient is under 8 years and treatment with steroids or methotrexate has proven ineffective or is not tolerated at a therapeutic dose; or disease requires control to prevent irreversible vision loss prior to achieving a therapeutic dose of methotrexate

## $\label{eq:Renewal} \textbf{Renewal} - \textbf{chronic ocular inflammation}$

Current approval Number (if known): Applications from any relevant practitioner. Approvals valid for 12 months. <b>Prerequisites</b> (tick boxes where appropriate)	
The patient has had a good clinical response following 3 initial doses	
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 uver cystoid macular oedema)	eitic
Following each 12 month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 1 daily, or steroid drops less than twice daily if under 18 years old	0mg
Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely high risk of irreversi vision loss if infliximab is withdrawn.	ole

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Infliximab - continued

Initial application — fistulising Crohn's disease Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)				
	and		Patient has confirmed Crohn's disease	
		or	Patient has one or more complex externally draining enterocutaneous fistula(e)	
		or	Patient has one or more rectovaginal fistula(e)	
			Patent has complex peri-anal fistula	
Rene	wal –	– fis	tulising Crohn's disease	
Applic	cation	s on	al Number (if known): ly from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 2 years. tick boxes where appropriate)	
		or	The number of open draining fistulae have decreased from baseline by at least 50%	
			There has been a marked reduction in drainage of all fistula(e) from baseline (in the case of adult patients, as demonstrated by a reduction in the Fistula Assessment score), together with less induration and patient reported pain	
	and [		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	
Appli	ication	ns or	i <b>on — neurosarcoidosis</b> Ily from a neurologist or Practitioner on the recommendation of a neurologist. Approvals valid for 18 months. tick boxes where appropriate)	
	and		Patient has been diagnosed with neurosarcoiosis by a multidisciplinary team	
	and_		Patient has CNS involvement	
	and		Patient has steroid-refractory disease	
		or	IV cyclophosphamide has been tried	
			Treatment with IV cyclophosphamide is clinically inappropriate	

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## APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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Infliximab - continued		
Renewal — neurosarcoidosis 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 bee	n 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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Fax Number:		Fax Number:

## Infliximab - continued

Applic	catior	is only	/ fror	<b>plaque psoriasis</b> n a dermatologist or any relevant practitioner on the recommendation of a dermatologist. Approvals valid for 3 months. xes where appropriate)
		[ and		The patient has had an initial Special Authority approval for adalimumab, etanercept or secukinumab for severe chronic plaque psoriasis
			or	Patient has experienced intolerable side effects from adalimumab, etanercept or secukinumab
				Patient has received insufficient benefit from adalimumab, etanercept or secukinumab to meet the renewal criteria for adalimumab, etanercept or secukinumab for severe chronic plaque psoriasis
	or			
			or or	<ul> <li>Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis</li> <li>Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis</li> <li>Patient has severe chronic localised genital or flexural plaque psoriasis where the plaques or lesions have been present for at least 6 months from the time of initial diagnosis.</li> <li>Patient has severe chronic localised genital or flexural plaque psoriasis where the plaques or lesions have been present for at least 6 months from the time of initial diagnosis, and with a Dermatology Life Quality Index (DLQI) score greater than 10</li> </ul>
		and [ and		Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin
		and		A PASI assessment has been completed for at least the most recent prior treatment course (but preferably all prior treatment courses), preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course
		[		The most recent PASI assessment is no more than 1 month old at the time of initiation
while s face, h severe	still o nand, e, and	n treat foot, g I for th	tmen genit ne fa	sponse" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably it but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the ital or flexural areas at least 2 of the 3 PASI symptom subscores for erythema, thickness and scaling are rated as severe or very ce, palm of a hand or sole of a foot the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed n treatment but no longer than 1 month following cessation of the most recent prior treatment.

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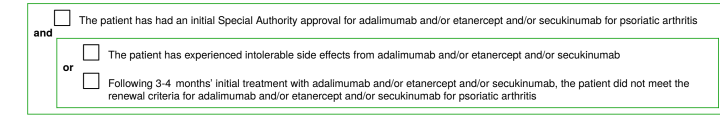
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Infliximat	<b>)</b> - co	ntinued			
Renewal –	– plac	ue psoria	sis		
Application	s from	any releva	,	vals valid for 6 months.	]
	or	and and or	Following each prior ir sustained at this level Patient had severe ch Following each all 3 of erythem course baseline Following each	dy" severe chronic plaque psoriasis at the start of tre infliximab treatment course the patient has a PASI sco when compared with the pre-infliximab treatment ba ronic plaque psoriasis of the face, or palm of a hand prior infliximab treatment course the patient has a rea a, thickness and scaling, to slight or better, or sustain a values prior infliximab treatment course the patient has a rea tained at this level, as compared to the pre-infliximab	ore which is reduced by 75% or more, or is seline value or sole of a foot at the start of treatment duction in the PASI symptom subscores for ed at this level, as compared to the treatment duction of 75% or more in the skin area
	or	and	The patient has compared to the	ronic localised genital or flexural plaque psoriasis at t experienced a reduction of 75% or more in the skin a e pre-treatment baseline value ermatology Quality of Life Index (DLQI) improvement ncing infliximab	area affected, or sustained at this level, as
and [	Iı	nfliximab to	be administered at do	ses no greater than 5 mg/kg every 8 weeks	

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### Infliximab - continued

itial application — previous use pplications from any relevant practitioner. Approvals valid for 6 months. rerequisites(tick boxes where appropriate)				
and	Patient was being treated with infliximab prior to 1 February 2019			
0	Rheumatoid arthritis			
0	Ankylosing spondylitis			
0	Psoriatic arthritis			
0	Severe ocular inflammation			
0	Chronic ocular inflammation			
0	Crohn's disease (adults)			
0	Crohn's disease (children)			
0	Fistulising Crohn's disease			
0	Severe fulminant ulcerative colitis			
0	Severe ulcerative colitis			
	Plaque psoriasis			
0	Neurosarcoidosis			
0	Severe Behcet's disease			
olications of	ntion — psoriatic arthritis only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 4 months. s(tick boxes where appropriate)			



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	ab - continued
Renewa	I — psoriatic arthritis
	approval Number (if known):
	ions only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. iisites(tick boxes where appropriate)
	<ul> <li>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</li> <li>The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior infliximab treatment in the opinion of the treating physician</li> </ul>
ar	Infliximab to be administered at doses no greater than 5 mg/kg every 8 weeks
Applicat	pplication — rheumatoid arthritis tions only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 4 months. isites(tick boxes where appropriate)
ar	The patient has had an initial Special Authority approval for adalimumab and/or etanercept for rheumatoid arthritis
	The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept or
	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
ar	Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance
Renewa	I — rheumatoid arthritis
Current	approval Number (if known):
Applicati	ions only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. iisites(tick boxes where appropriate)
ar	Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance
	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 treatment in the opinion of the physician
ar	nd Infliximab to be administered at doses no greater than 3 mg/kg every 8 weeks

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## 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) and				
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; azathioprine, steroids, 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. Approvals valid for 6 months.				
Prerequisites(tick boxes where appropriate)				
Patient has had a good clinical response to initial treatment with measurably improved quality of life and				
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. Approvals valid for 2 years.				
Prerequisites(tick boxes where appropriate)				
Where maintenance treatment is considered appropriate, infliximab should be used in combination with immunomodulators and reassessed every 6 months and				
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				

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### Infliximab - continued

Appl	icatior	ication — severe ocular inflammation ns from any relevant practitioner. Approvals valid for 4 months. tes(tick boxes where appropriate)	
		and The patient has had an initial Special Authority approval for adalimumab for severe ocular inflammation 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 severe ocular inflammation	
	or	and Patient has severe, vision-threatening ocular inflammation requiring rapid control and Patient has severe, vision-threatening ocular inflammation requiring rapid control I I I I I I I I I I I I I I I I I I I	
		ineffective at controlling symptoms	

Renewal — severe ocular inflammation
Current approval Number (if known):
Applications from any relevant practitioner. Approvals valid for 12 months. <b>Prerequisites</b> (tick boxes where appropriate)
The patient has had a good clinical response following 3 initial doses     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 uveitic
or cystoid macular oedema) Following each 12 month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10mg 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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Fax Number:		Fax Number:

Infliximab - continued

Initial application — ulcerative colitis Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)	
Patient has active ulcerative colitis	
or Patients SCCAI is greater than or equal to 4	
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 t immunomodulators and systemic corticosteroids	herapy with
Renewal — ulcerative colitis	
Current approval Number (if known):	
Applications from any relevant practitioner. Approvals valid for 2 years.	
Prerequisites(tick boxes where appropriate)	
or The SCCAI score has reduced by 2 points or more from the SCCAI score when the patient was initiated on inflixing The PUCAI score has reduced by 10 points or more from the PUCAI score when the patient was initiated on inflixing the public of the PUCAI score has reduced by 10 points or more from the PUCAI score when the patient was initiated on inflixing the public of the publ	
and 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 to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered after completing the last re-induction cycle. Up to 10 mg/kg every 8 weeks (or equivalent) may be used for patients treate prior to 1 February 2019	sixteen weeks
Initial application — pyoderma gangrenosum Applications only from a dermatologist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)	
Patient has pyoderma gangrenosum* and	
Patient has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisor azathioprine, or methotrexate) and not received an adequate response and	ne, ciclosporine,
A maximum of 8 doses	

Note: Note: Indications marked with \* are unapproved indications.

Enquiries	to	Ministry	of	Health
0800 855	06	6		

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Fax Number:		Fax Number:			
Infliximab - continued					
Renewal — pyoderma gangrenosum					
Current approval Number (if known):					
	ner on the recommendation of a dermatologist. Appr	ovals valid for 4 months.			
Prerequisites(tick boxes where appropriate)					
Patient has shown clinical improve	ement				
Patient continues to require treatm	ent				
A maximum of 8 doses					
Initial application — inflammatory bowel arthri Applications from any relevant practitioner. Appro					
Prerequisites(tick boxes where appropriate)					
	lcerative colitis or active Crohn's disease				
And Patient has had axial inflammatory	pain for six months or more				
and Patient is unable to take NSAIDs					
and Patient has unequivocal sacroiliitis demonstrated by radiological imaging or MRI					
and					
<ul> <li>Patient's disease has not responded adequately to prior treatment consisting of at least 3 months of an exercise regime supervised by a physiotherapist</li> <li>and</li> </ul>					
	on a 0 - 10 scale completed after the 3 month exe	rcise trial, but prior to ceasing any previous			
Renewal — inflammatory bowel arthritis – axial					
Current approval Number (if known): Applications from any relevant practitioner. Approv					

**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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### Infliximab - continued

olications	ation — inflammatory bowel arthritis – peripheral from any relevant practitioner. Approvals valid for 6 months. s(tick boxes where appropriate)		
and	<ul> <li>Patient has a diagnosis of active ulcerative colitis or active Crohn's disease</li> <li>Patient has active arthritis in at least four joints from the following: hip, knee, ankle, subtalar, tarsus, forefoot, wrist, elbow, shoulder, sternoclavicular</li> </ul>		
and			
and	Patient has tried and not experienced a response to at least three months of methotrexate or azathioprine at a maximum tolerated dose (unless contraindicated)		
and	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 or		
	Patient has an ESR greater than 25 mm per hour measured no more than one month prior to the date of this application or		
	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		
newal — i	inflammatory bowel arthritis – peripheral		
rent appro	oval Number (if known):		
lications	from any relevant practitioner. Approvals valid for 2 years.		

Following initial treatment, patient has experienced at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician

Patient has experienced at least a continuing 30% improvement in active joint count from baseline in the opinion of the treating physician

I confirm the above details are correct and that in signing this form I understand I may be audited.

or