SA2157 - Adalimumab (Humira - Alternative brand)

Arthritis - polyarticular course juvenile idiopathic - Initial application10
Arthritis - polyarticular course juvenile idiopathic - Renewal
Arthritis - psoriatic - Initial application11
Arthritis - psoriatic - Renewal
Arthritis – oligoarticular course juvenile idiopathic - Initial application10
Arthritis – oligoarticular course juvenile idiopathic - Renewal
Arthritis - rheumatoid - Initial application11
Arthritis – rheumatoid - Renewal
Behcet's disease – severe - Initial application2
Behcet's disease - severe - Renewal2
Crohn's disease - adult - Initial application
Crohn's disease - adult - Renewal
Crohn's disease - children - Initial application6
Crohn's disease - children - Renewal
Crohn's disease - fistulising - Initial application
Crohn's disease - fistulising - Renewal
Hidradenitis suppurativa - Initial application
Hidradenitis suppurativa - Renewal
Ocular inflammation – chronic - Initial application
Ocular inflammation – chronic - Renewal
Ocular inflammation – severe - Initial application
Ocular inflammation – severe - Renewal
Psoriasis - severe chronic plaque - Initial application
Psoriasis - severe chronic plaque - Renewal
Pyoderma gangrenosum - Initial application
Pyoderma gangrenosum - Renewal5
Still's disease – adult-onset (AOSD) - Initial application
Still's disease – adult-onset (AOSD) - Renewal
Ankylosing spondylitis - Initial application9
Ankylosing spondylitis - Renewal

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 (Humira - Alternative brand)

Initial application — Behcet's disease – severe Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
or The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen			
 and Patient has received a maximum of 6 months treatment with Amgevita and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and Adalimumab to be administered at doses no greater than 40 mg every 14 days 			
Renewal — Behcet's disease – severe Current approval Number (if known): Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
Image: The patient has had a good clinical response to treatment with measurably improved quality of life and Image: Adalimumab to be administered at doses no greater than 40 mg every 14 days			
Initial application — Hidradenitis suppurativa Applications only from a dermatologist or Practitioner on the recommendation of a dermatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
or Patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen			
and Patient has received a maximum of 6 months treatment with Amgevita and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and			
Adalimumab to be administered at doses no greater than 40 mg every 7 days. Fortnightly dosing has been considered			

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 (Humira - Alternative brand) - continued

Renewal — Hidradenitis suppurativa				
Current approval Number (if known):				
Applications only from a dermatologist or Practitioner on the recommendation of a dermatologist. Approvals valid for 6 months. Prerequisites (tick boxes where appropriate)				
The patient has a reduction in active lesions (e.g. inflammatory nodules, abscesses, draining fistulae) of 25% or more from baseline and				
The patient has a Dermatology Quality of Life Index improvement of 4 or more from baseline and				
Adalimumab is to be administered at doses no greater than 40mg every 7 days. Fortnightly dosing has been considered				
Initial application — Psoriasis - severe chronic plaque Applications only from a dermatologist or Practitioner on the recommendation of a dermatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)				
The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment or				
Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen				
and Patient has received a maximum of 6 months treatment with Amgevita				
and				
and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and				

Enquiries to Ministry of Health
0800 855 066

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 (Humira - Alternative brand) - continued

			riasis - severe chronic plaque Number (if known):				
Applic	cations	s only	r from a dermatologist or Practitioner on the recommendation of a dermatologist. Approvals valid for 6 months. ck boxes where appropriate)				
			Patient had "whole body" severe chronic plaque psoriasis at the start of treatment and Following each prior adalimumab treatment course the patient has a PASI score which is reduced by 75% or more, or is sustained at this level, when compared with the pre-adalimumab treatment baseline value or Following each prior adalimumab treatment course the patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, when compared with the pre-treatment baseline value				
		or	Patient had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment and Following each prior adalimumab treatment course the patient has a reduction in the PASI symptom subscores for all 3 of erythema, thickness and scaling, to slight or better, or sustained at this level, as compared to the treatment course baseline values Following each prior adalimumab treatment course the patient has a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-adalimumab treatment baseline value				
	and Adalimumab to be administered at doses no greater than 40 mg every 14 days						
Appli	nitial application — Pyoderma gangrenosum Applications only from a dermatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)						
		or	 The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen 				
	and [F	Patient has received a maximum of 6 months treatment with Amgevita				
	Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and						
		A maximum of 8 doses					

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 (Humira - Alternative brand) - continued

Renewal — Pyoderma gangrenosum			
Current approval Number (if known): Applications only from a dermatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
The patient has demonstrated clinical improvement and continues to require treatment A maximum of 8 doses			
Initial application — Crohn's disease - adult Applications only from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
 The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevitat Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita and clinician attributes this loss of disease response to a change in treatment regimen Patient has Crohn's and is considered to be at risk of disease destabilisation if there were to be a change to current treatment 			
and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and Adalimumab to be administered at doses no greater than 40 mg every 14 days			

Renewal — Crohn's disease - adult

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

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

	CDAI score has reduced by 100 points from the CDAI score when the	patient was initiated on adalimumab
	CDAI score is 150 or less	
	The patient has demonstrated an adequate response to treatment, but	CDAI score cannot be assessed
and [and Adalimumab to be administered at doses no greater than 40 mg every 14 day	/s

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

Signed: Date:

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 (Humira - Alternative brand) - continued

Initial application — Crohn's disease - children Applications only from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)
 The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita and clinician attributes this loss of disease response to a change in treatment regimen Patient has Crohn's and is considered to be at risk of disease destabilisation if there were to be a change to current treatment Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication Adalimumab to be administered at doses no greater than 40 mg every 14 days
Renewal — Crohn's disease - children Current approval Number (if known): Applications only from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 6 months.

Prerequisites(tick boxes where appropriate)

or	PCDAI score has reduced by 10 points from the PCDAI score when the patient was initiated on adalimumab			
	PCDAI score is 15 or less			

The patient has demonstrated an adequate response to treatment, but PCDAI score cannot be assessed

Adalimumab to be administered at doses no greater than 40 mg every 14 days

Initial application — Crohn's disease - fistulising

and

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

	[or	The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita
	5" [pr [Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita and clinician attributes this loss of disease response to a change in treatment regimen Patient has Crohn's and is considered to be at risk of disease destabilisation if there were to be a change to current treatment
and	-	atient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication dalimumab to be administered at doses no greater than 40 mg every 14 days

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 (Humira - Alternative brand) - continued

Renewal — Crohn's disease - fistulising			
Current approval Number (if known):			
Applications only from a gastroenterologist or Practitioner on the recommendation of a gastroenterologist. Approvals valid for 6 months. Prerequisites (tick boxes where appropriate)			
The number of open draining fistulae have decreased from baseline by at least 50% or There has been a marked reduction in drainage of all fistula(e) from baseline as demonstrated by a reduction in the Fistula			
Assessment score, together with less induration and patient-reported pain			
and Adalimumab to be administered at doses no greater than 40 mg every 14 days			
Initial application — Ocular inflammation – chronic Applications from any relevant practitioner. Approvals valid for 12 months. Prerequisites(tick boxes where appropriate)			
The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita			
Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with Amgevita, and a maximum of 6 months treatment with Amgevita and clinician attributes this loss of disease response to a change in treatment regimen			
or Patient has uveitis and is considered to be at risk of vision loss if they were to change treatment			
and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and			
Adalimumab to be administered at doses no greater than 40 mg every 14 days			

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 (Humira - Alternative brand) - continued

Renewal — Ocular inflammation – chronic			
Current approval Number (if known):			
Applications from any relevant practitioner. Approvals valid for 12 months. Prerequisites (tick boxes where appropriate)			
 The patient has had a good clinical response following 12 weeks' initial treatment 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 uveitic cystoid macular oedema) or 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 			
Initial application — Ocular inflammation – severe Applications from any relevant practitioner. Approvals valid for 12 months. Prerequisites(tick boxes where appropriate)			
 The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment, and a maximum of 6 months treatment with Amgevita Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with Amgevita, and a maximum of 6 months treatment with Amgevita and clinician attributes this loss of disease response to a change in treatment regimen Patient has uveitis and is considered to be at risk of vision loss if they were to change treatment 			
and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and Adalimumab to be administered at doses no greater than 40 mg every 14 days			

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 (Humira - Alternative brand) - continued

Renewal — Ocular inflammation – severe			
Current approval Number (if known):			
Applications from any relevant practitioner. Approvals valid for 12 months. Prerequisites (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 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 Adalimumab to be administered at doses no greater than 40 mg every 14 days 			
Initial application — ankylosing spondylitis Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
or Patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita)			
and Patient has received a maximum of 6 months treatment with Amgevita and			
Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and			
Adalimumab to be administered at doses no greater than 40 mg every 14 days			
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)

> 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

Adalimumab to be administered at doses no greater than 40 mg every 14 days

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

and

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 (Humira - Alternative brand) - continued

Initial application — Arthritis – oligoarticular course juvenile idiopathic Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
or The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen			
and Patient has received a maximum of 6 months treatment with Amgevita and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication			
Renewal — Arthritis – oligoarticular course juvenile idiopathic			
Current approval Number (if known): Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites (tick box where appropriate)			
The patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline			
Initial application — Arthritis - polyarticular course juvenile idiopathic Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
or Patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen			
and Patient has received a maximum of 6 months treatment with Amgevita and			
Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication			
Renewal — Arthritis - polyarticular course juvenile idiopathic			
Current approval Number (if known): Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites (tick box where appropriate)			
The patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline			
I confirm the above details are correct and that in signing this form I understand I may be audited.			

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 (Humira - Alternative brand) - continued

Initial application — Arthritis - psoriatic

Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites (tick boxes where appropriate)
The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment or
Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen
and Patient has received a maximum of 6 months treatment with Amgevita and
Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and
Adalimumab to be administered at doses no greater than 40 mg every 14 days
Renewal — Arthritis - psoriatic
Current approval Number (if known): Applications only from a named specialist, rheumatologist or Practitioner on the recommendation of a named specialist or rheumatologist. Approvals valid for 6 months. Prerequisites (tick boxes where appropriate)
The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior adalimumab treatment in the opinion of the treating physician Adalimumab to be administered at doses no greater than 40 mg every 14 days
Initial application — Arthritis – rheumatoid Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)
The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment or
Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen
and Patient has received a maximum of 6 months treatment with Amgevita and
Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication and
Adalimumab to be administered at doses no greater than 40 mg every 14 days
Patient cannot take concomitant methotrexate and requires doses of adalimumab higher than 40 mg every 14 days to maintain an adequate response

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 (Humira - Alternative brand) - continued

Renewal — Arthritis – rheumatoid			
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)			
The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior adalimumab treatment in the opinion of the treating physician and			
Adalimumab to be administered at doses no greater than 40 mg every 14 days			
Patient cannot take concomitant methotrexate and requires doses of adalimumab higher than 40 mg every 14 days to maintain an adequate response			
Initial application — Still's disease – adult-onset (AOSD) Applications only from a rheumatologist or Practitioner on the recommendation of a rheumatologist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)			
The patient has experienced intolerable side effects from adalimumab (Amgevita) following a minimum of 4 weeks treatment or			
Patient has developed symptoms of loss of disease control following a minimum of 4 weeks treatment with adalimumab (Amgevita) and clinician attributes this loss of disease response to a change in treatment regimen			
and Patient has received a maximum of 6 months treatment with Amgevita			
and Patient has previously had a Special Authority approval for the Humira brand of adalimumab for this indication			
Renewal — Still's disease – adult-onset (AOSD)			
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 box where appropriate)			

The patient has demonstrated a sustained improvement in inflammatory markers and functional status