RS2063 - Adalimumab (Amgevita)

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Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:

Adalimumab (Amgevita)

INITIATION – Behcet's disease - severe		
Prerequisites (tick boxes where appropriate)		

O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.

ana	($\overline{)}$	Tho	patient has severe Behcet's disease* that is significantly impacting the patient's quality of life	
	and				
			0	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)	
		or	0	The patient has severe gastrointestinal, rheumatological and/or mucocutaneous symptoms and has not responded adequately to two or more treatments appropriate for the particular symptom(s)	
Note:	lote: Indications marked with * are unapproved indications.				

INITIATION – Hidradenitis suppurativa Re-assessment required after 4 months

Prerequisites (tick boxes where appropriate)

O Prescribed by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.						
	O Patient has hidradenitis suppurativa Hurley Stage II or Hurley Stage III lesions in distinct anatomic areas					
	and	Patient has tried, but had an inadequate response to at least a 90 day trial of systemic antibiotics or patient has demonstrated intolerance to or has contraindications for systemic antibiotics				
	and O Patient has 3 or more active lesions					
		The patient has a DLQI of 10 or more and the assessment is no more than 1 month old at time of application				

TINUATION – Hidradenitis suppurativa ssessment required after 2 years
equisites (tick boxes where appropriate)
Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.
O The patient has a reduction in active lesions (e.g. inflammatory nodules, abscesses, draining fistulae) of 25% or more from baseline and O The patient has a DLQI improvement of 4 or more from baseline

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I confirm that the above details are correct:

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PRES	PRESCRIBER			PATIENT:
Name	Name:			Name:
Ward	:			NHI:
Ada	limu	mab	(Am	gevita) - continued
Re-a	assess	sment i	equi	psoriasis - severe chronic ed after 4 months ixes where appropriate)
O Prescribed by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endors Hospital.			by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ	
		(and	С	Patient has had an initial Special Authority approval for etanercept for severe chronic plaque psoriasis
	$ \begin{array}{ c c c } O & \text{Patient has experienced intolerable side effects} \\ \text{or} & \\ O & Patient has received insufficient benefit to meet the set of the s$		or	 Patient has experienced intolerable side effects Patient has received insufficient benefit to meet the renewal criteria for etanercept for severe chronic plaque psoriasis
	or			
or				
			or	O 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
				O 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
following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or a		Patient has tried, but had an inadequate response 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 (С	A PASI assessment or (DLQI) assessment has been completed for at least the most recent prior treatment course but no longer than 1 month following cessation of each prior treatment course and is no more than 1 month old at the time of application

I confirm that the above details are correct:

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PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:
Adalimumab (Amgevita) - continued	
CONTINUATION – Plaque psoriasis - severe chronic Re-assessment required after 2 years	
Prerequisites (tick boxes where appropriate)	
O Patient had "whole body" severe chronic plaque psorias	sis at the start of treatment
O The patient has experienced a 75% or more redu	ction in PASI score, or is sustained at this level, when compared with

		or ()	the pre-treatment baseline value The patient has a DLQI improvement of 5 or more, when compared with the pre-treatment baseline value
or) Patie	ent had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment
	and	or O	The patient has experienced 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 The patient has experienced a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-treatment baseline value
or	and) Patie	ent had severe chronic localised genital or flexural plaque psoriasis at the start of treatment
		or O	The patient has experienced a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-treatment baseline value Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing adalimumab

		pyoderma gangrenosum (tick boxes where appropriate)
and	Pres Hosp	cribed by, or recommended by a dermatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ ital.
an		Patient has pyoderma gangrenosum* Patient has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclosporin, azathioprine, or methotrexate) and not received an adequate response
Note: In	dicatio	ons marked with * are unapproved indications.

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PRES	SCRIE	BER		PATIENT:	
Name:				Name:	
Ward	:			NHI:	
Adal	imu	mał	b (Amgevita) - continued		
Re-a	ssess equis	smer sites Preso	Crohn's disease - adults nt required after 6 months (tick boxes where appropriate) cribed by, or recommended by any relevant practitioner, or in acc lospital.	cordance with a protocol or guideline that has been endorsed by the Health	
	and	0	Patient has severe active Crohn's disease		
		or	\sim		
		or	 Patient has extensive small intestine disease affecting mo Patient has evidence of short gut syndrome or would be a 		
		or	O Patient has an ileostomy or colostomy and has intestinal	inflammation	
	and	0	Patient has tried but had an inadequate response to, or has exp and corticosteroids	perienced intolerable side effects from, prior therapy with immunomodulators	
	equis	sites Preso	nt required after 2 years (tick boxes where appropriate) cribed by, or recommended by any relevant practitioner, or in acc lospital.	cordance with a protocol or guideline that has been endorsed by the Health	
	or	0	CDAI score has reduced by 100 points from the CDAI score, or adalimumab	HBI score has reduced 3 points, from when the patient was initiated on	
	-	Ο	CDAI score is 150 or less, or HBI is 4 or less		
	or	0	The patient has demonstrated an adequate response to treatme	ent, but CDAI score and/or HBI score cannot be assessed	
Re-a	ssess equis	smer sites Preso	Crohn's disease - children nt required after 6 months (tick boxes where appropriate) cribed by, or recommended by any relevant practitioner, or in acc dospital.	cordance with a protocol or guideline that has been endorsed by the Health	
	and	Ο	Paediatric patient has active Crohn's disease		
		or	 Patient has a PCDAI score of greater than or equal to 30 Patient has extensive small intestine disease 		
	and			perienced intolerable side effects from, prior therapy with immunomodulators	

Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRES	SCRIBE		PATIENT:
Name:			Name:
Ward	:		NHI:
Ada	limum	b (Amgevita) - continued	
Re-a	issessm	ON – Crohn's disease - children nt required after 2 years (tick boxes where appropriate)	
and		cribed by, or recommended by any relevant practitioner Hospital.	r, or in accordance with a protocol or guideline that has been endorsed by the Health
	or	PCDAI score has reduced by 10 points from the PCDA	Al score when the patient was initiated on adalimumab
	or	PCDAI score is 15 or less	
		The patient has demonstrated an adequate response	to treatment but PCDAI score cannot be assessed
Re-a	equisit	Hospital. Patient has confirmed Crohn's disease O Patient has one or more complex externally draited O Patient has one or more rectovaginal fistula(e) O Patient has complex peri-anal fistula	r, or in accordance with a protocol or guideline that has been endorsed by the Health ining enterocutaneous fistula(e) and is no more than 1 month old at the time of application
CONTINUATION – Crohn's disease - fistulising Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the H NZ Hospital.			
	or	The number of open draining fistulae have decreased There has been a marked reduction in drainage of all score, together with less induration and patient-report	fistula(e) from baseline as demonstrated by a reduction in the Fistula Assessment

or

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRES	SCRIE	BER		PATIENT:		
Name	e:					
Ward	:			NHI:		
Ada	limu	mab	(An	ngevita) - continued		
				ur inflammation - chronic uired after 4 months		
				poxes where appropriate)		
and		Presci NZ Ho		by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health al.		
	or	O	The p	patient has had an initial Special Authority approval for infliximab for chronic ocular inflammation		
		and	O or or	O Patient is under 18 years and treatment with methotrexate has proven ineffective or is not tolerated at a therapeutic dose		
				O 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		
	\square	_				
Re-a	issess equis	sment sites (requ tick b ribed	Dcular inflammation - chronic uired after 2 years poxes where appropriate) I by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health al.		
	O The patient has had a good clinical response following 12 weeks' initial treatment or O Following each 0, were treatment against the restingt has had a sustained reduction in inflammation (Chardendisation of Llupikia					

Following each 2 year 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 2 year 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

Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRES	SCRIE	BER	PATIENT:			
Name	ə:					
Ward	:		NHI:			
Adal	limu	mab	b (Amgevita) - continued			
INITIATION – Ocular inflammation - severe Re-assessment required after 4 months Prerequisites (tick boxes where appropriate)						
(and	C	Presci	cribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health lospital.			
and	or	0	Patient has had an initial Special Authority approval for infliximab for severe ocular inflammation			
		and	O Patient has severe, vision-threatening ocular inflammation requiring rapid control			
			O Treatment with high-dose steroids (intravenous methylprednisolone) followed by high dose oral steroids has proven ineffective at controlling symptoms O Patient developed new inflammatory symptoms while receiving high dose steroids			
			O Patient is aged under 8 years and treatment with high dose oral steroids and other immunosuppressants has proven ineffective at controlling symptoms			
Re-a	CONTINUATION – Ocular inflammation - severe Re-assessment required after 2 years Prerequisites (tick boxes where appropriate)					
O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorse NZ Hospital.						
	or	0	The patient has had a good clinical response following 3 initial doses			
	or		Following each 2 year treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < $\frac{1}{2}$ + anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uveitic cystoid macular oedema)			

O Following each 2 year 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

Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRESCRIE	BER	PATIENT:
Name:		
Ward:		NHI:
Adalimu	mab (Ar	ngevita) - continued
Re-assess Prerequis	sment requ sites (tick b	besing spondylitis ired after 6 months ioxes where appropriate) by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
	and	 Patient has had an initial Special Authority approval for etanercept for ankylosing spondylitis O The patient has experienced intolerable side effects O The patient has received insufficient benefit to meet the renewal criteria for ankylosing spondylitis
or Patient has a confirmed diagnosis of ankylosing spondylitis for m and Patient has low back pain and stiffness that is relieved by exercise and Patient has bilateral sacroiliitis demonstrated by radiology imagin and Patient has not responded adequately to treatment with two or m		Patient has a confirmed diagnosis of ankylosing spondylitis for more than six months Patient has low back pain and stiffness that is relieved by exercise but not by rest Patient has bilateral sacroiliitis demonstrated by radiology imaging Patient has not responded adequately to treatment with two or more NSAIDs, while patient was undergoing at least 3 months of a regular exercise regimen for ankylosing spondylitis
	and or and O	 Patient has limitation of motion of the lumbar spine in the sagittal and the frontal planes as determined by the following BASMI measures: a modified Schober's test of less than or equal to 4 cm and lumbar side flexion measurement of less than or equal to 10 cm (mean of left and right) Patient has limitation of chest expansion by at least 2.5 cm below the average normal values corrected for age and gender A BASDAI of at least 6 on a 0-10 scale completed after the 3 month exercise trial, but prior to ceasing any previous pharmacological treatment and is no more than 1 month old at the time of application
Re-assess Prerequis	sment requ sites (tick b	nkylosing spondylitis ired after 2 years iox where appropriate) by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health

NZ Hospital.

For applications where 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

and

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PRE	SCRIE	BER	PATIENT:				
Name	ə:		Name:				
Ward	:		NHI:				
Ada	dalimumab (Amgevita) - continued						
INIT Re-a	 At least 2 active joints with limited range of motion, pain or tenderness after a 3-month trial of methotrexate (at the maximum tolerated dose) 						
CONTINUATION – Arthritis - oligoarticular course juvenile idiopathic Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed							
and	or	0	ospital. Following initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline				

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Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

PRESCRIBER	PATIENT:			
Name:	Name:			
Ward:	NHI:			
Adalimumab (Amgevita) - continued				
INITIATION – Arthritis - polyarticular course juvenile idiopathic Re-assessment required after 6 months Prerequisites (tick boxes where appropriate)	logist, or in accordance with a protocol or guideline that has been endorsed			
O Patient has had an initial Special Authority approval for and O Patient has experienced intolerable side effects or O Patient has received insufficient benefit to meet th	etanercept for polyarticular course juvenile idiopathic arthritis (JIA) ne renewal criteria for polyarticular course JIA			
and O Patient has had polyarticular course JIA for 6 months du and O At least 5 active joints and at least 3 joints with li or or	mited range of motion, pain or tenderness after a 3-month trial of ore of at least 2.5) after a 3-month trial of methotrexate (at the			
CONTINUATION – Arthritis - polyarticular course juvenile idiopathic Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the He NZ Hospital.				

O Following initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline

On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline

or

PRES	CRIB	ER		PATIENT:
Name	:			Name:
Ward:				NHI:
Adal	imur	nab	(An	ngevita) - continued
Re-a	ssess equisi	iment r ites (ti	requi ck b bed	ris - psoriatic ired after 6 months oxes where appropriate) by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ
and		(and	Cor	Patient has had an initial Special Authority approval for etanercept or secukinumab for psoriatic arthritis O Patient has experienced intolerable side effects O Patient has received insufficient benefit to meet the renewal criteria for psoriatic arthritis
	or	and (and (and	O O O or or	 Patient has had active psoriatic arthritis for six months duration or longer Patient has tried and not responded to at least three months of methotrexate at a maximum tolerated dose (unless contraindicated) Patient has tried and not responded to at least three months of sulfasalazine or leflunomide at maximum tolerated doses (unless contraindicated) Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen joints Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip Patient has CRP level greater than 15 mg/L measured no more than one month prior to the date of this application Patient has an elevated ESR greater than 25 mm per hour ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day
Re-a Prere	ssess equisi	iment r ites (ti	requi ick b bed	and has done so for more than three months rthritis - psoriatic ired after 2 years oxes where appropriate) by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health I.
and	or (re O F	espo Patier	wing initial treatment, the patient has at least a 50% decrease in swollen joint count from baseline and a clinically significant onse in the opinion of the physician and demonstrates at least a continuing 30% improvement in swollen joint count from baseline and a clinically significant response opinion of the treating physician

PRESC	CRIBI	ER		PATIENT:		
Name:				Name:		
Ward:				NHI:		
Adalir	mun	nab	(An	ngevita) - continued		
INITIA Re-ass Prerect	ATION sessr quisi P	Hospital.		is - rheumatoid ired after 6 months oxes where appropriate) by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ The patient has had an initial Special Authority approval for etanercept for rheumatoid arthritis O The patient has experienced intolerable side effects O The patient has received insufficient benefit from etanercept to meet the renewal criteria for rheumatoid arthritis Patient has had rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity		
		and (and and	or or	 Patient has tried and not responded to at least three months of methotrexate in combination with sulfasalazine and hydroxychloroquine sulphate at maximum tolerated doses (unless contraindicated) Patient has tried and not responded to at least three months of methotrexate in combination with the maximum tolerated dose of ciclosporin Patient has tried and not responded to at least three months of therapy at the maximum tolerated dose of leflunomide alone or in combination with methotrexate Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen joints Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip 		
Re-ass	sessr quisi t	nent r tes (ti	equi ck b	rthritis - rheumatoid ired after 2 years oxes where appropriate) by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health		

and		NZ F	ospital.
	or	0	Following 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
		0	On subsequent reapplications, 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

PRES	SCRI	BER	PATIENT:
Name	e:		
Ward			NHI:
Adal	imu	ımal	o (Amgevita) - continued
			Still's disease - adult-onset (AOSD) (tick boxes where appropriate)
	- C		
and		Hosp	cribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ ital.
unu		an	O The patient has had an initial Special Authority approval for etanercept and/or tocilizumab for (AOSD)
			O Patient has experienced intolerable side effects from etanercept and/or tocilizumab
			or O Patient has received insufficient benefit from at least a three-month trial of etanercept and/or tocilizumab
	or		
		an	O Patient diagnosed with AOSD according to the Yamaguchi criteria
		an	O 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
			O Patient has persistent symptoms of disabling poorly controlled and active disease
Re-a Prero	sses	smer sites Pres	ulcerative colitis It required after 6 months (tick boxes where appropriate) cribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health lospital.
and		O	Patient has active ulcerative colitis
	and		O Patient's SCCAI score is greater than or equal to 4
		or	O Patient's PUCAI score is greater than or equal to 20
	and and	0	Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior therapy with immunomodulators and systemic corticosteroids
l		0	Surgery (or further surgery) is considered to be clinically inappropriate
Re-a	sses	smer sites Pres	DN – ulcerative colitis It required after 2 years (tick boxes where appropriate) cribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health
and			The SCCAL energy has reduced by 2 points or more from the SCCAL energy when the potient was initiated on historia therepy
	or	\bigcirc	The SCCAI score has reduced by 2 points or more from the SCCAI score when the patient was initiated on biologic therapy
		\cup	The PUCAI score has reduced by 10 points or more from the PUCAI score when the patient was initiated on biologic therapy

PRES	CRIBE	R PATIENT:			
Name	:	Name:			
Ward: NHI:					
Adal	imum	ab (Amgevita) - continued			
INITIATION – undifferentiated spondyloarthiritis Re-assessment required after 6 months Prerequisites (tick boxes where appropriate)					
and		escribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ spital.			
	C	Patient has undifferentiated peripheral spondyloarthritis* with active peripheral joint arthritis in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip			
	and _	Patient has tried and not responded to at least three months of each of methotrexate, sulphasalazine and leflunomide, at maximum tolerated doses (unless contraindicated)			
		O Patient has a CRP level greater than 15 mg/L measured no more than one month prior to the date of this application			
		O Patient has an ESR greater than 25 mm per hour measured no more than one month prior to the date of this application			
		O ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months			
Note	Indica	tions marked with * are unapproved indications.			
Re-a	CION – undifferentiated spondyloarthiritis ent required after 2 years es (tick boxes where appropriate) escribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health Hospital.				
	or	 Following 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 in the opinion of the treating physician			
Re-a	ssessm	- inflammatory bowel arthritis – axial ent required after 6 months es (tick boxes where appropriate)			
(and		escribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ spital.			
	and	Patient has a diagnosis of active ulcerative colitis or active Crohn's disease			
	and	Patient has axial inflammatory pain for six months or more			
	С	Patient is unable to take NSAIDs			
	and	Patient has unequivocal sacroiliitis demonstrated by radiological imaging or MRI			
	and	Patient has not responded adequately to prior treatment consisting of at least 3 months of an exercise regime supervised by a physiotherapist			
	and	A BASDAI of at least 6 on a 0-10 scale completed after the 3 month exercise trial, but prior to ceasing any previous pharmacological treatment			

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PRESC	CRIBEF	ł	PATIENT:			
Name:			Name:			
Ward:			NHI:			
Adalii	muma	b (Amgevita) - continued				
		ION – inflammatory bowel arthritis – axial ent required after 2 years				
		s (tick box where appropriate)				
and		scribed by, or recommended by any relevant practitioner, or in Hospital.	accordance with a protocol or guideline that has been endorsed by the Health			
C		ere treatment has resulted in an improvement in BASDAI of 4 rovement in BASDAI of 50%, whichever is less	or more points from pre-treatment baseline on a 10 point scale, or an			
Re-as	sessme quisite:) Pre:	 N - inflammatory bowel arthritis - peripheral sment required after 6 months sites (tick boxes where appropriate) Prescribed by, or recommended by a rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital. O Patient has a diagnosis of active ulcerative colitis or active Crohn's disease 				
;	and and O and	sternoclavicular	owing: hip, knee, ankle, subtalar, tarsus, forefoot, wrist, elbow, shoulder, t three months of methotrexate, or azathioprine at a maximum tolerated			
	and _	Patient has tried and not experienced a response to at least contraindicated)	t three months of sulphasalazine at a maximum tolerated dose (unless			
	0	r O Patient has an ESR greater than 25 mm per hour	red no more than one month prior to the date of this application acceiving prednisone therapy at a dose of greater than 5 mg per day and			
Re-as	CONTINUATION – inflammatory bowel arthritis – peripheral Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by any relevant practitioner, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.					

Following 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

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

I confirm that the above details are correct:

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or ()