Enquiries to Ministry of Health 0800 855 066

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

Page 1 Form SA2183 April 2025

EANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:			
:	First Names:	First Names:			
	Surname:	Surname:			
S:	DOB:	Address:			
	Address:				
nber:		Fax Number:			
zumab					
Patient has active Crohn's disease Patient has had an initial app meet renewal criteria (unless or Patient has a CDAI score of g or Patient has extensive small in or Patient has evidence of short or	proval for prior biologic therapy and has experienced intolerable side effects or insufficient benefit to s contraindicated) greater than or equal to 300, or HBI score of greater than or equal to 10 intestine disease affecting more than 50 cm of the small intestine rt gut syndrome or would be at risk of short gut syndrome with further bowel resection				
or Patient has experienced intol	nomodulators and corticosteroids erable side effects from immunomodulators and cort				
Renewal — Crohn's disease - adults Current approval Number (if known):					
:	pplication — Crohn's disease - adults tions from any relevant practitioner. Approv uisites(tick boxes where appropriate) Patient has active Crohn's disease nd Patient has had an initial app meet renewal criteria (unless or Patient has extensive small in or Patient has evidence of shore or Patient has evidence of shore or Patient has an ileostomy or cond Patient has tried but experier from prior therapy with immu or Patient has experienced intol or Immunomodulators and cortical al — Crohn's disease - adults approval Number (if known):	First Names: Surname: DOB: Address: Address: Patient has active Crohn's disease - adults ions from any relevant practitioner. Approvals valid for 6 months. Jisites(tick boxes where appropriate) Patient has had an initial approval for prior biologic therapy and has experienced meet renewal criteria (unless contraindicated) Patient has a CDAI score of greater than or equal to 300, or HBI score of greater or Patient has extensive small intestine disease affecting more than 50 cm of the smort patient has evidence of short gut syndrome or would be at risk of short gut syndrome or would be at risk of short gut syndrome or would be at risk of short gut syndrome or would be at risk of short gut syndrome or patient has an ileostomy or colostomy, and has intestinal inflammation Patient has tried but experienced an inadequate response to (including lack of initing from prior therapy with immunomodulators and corticosteroids Patient has experienced intolerable side effects from immunomodulators and corticosteroids are contraindicated Patient has experienced intolerable side effects from immunomodulators and corticosteroids are contraindicated Cohi's disease - adults approval Number (if known):			

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Reg No	o:		First Names:	First Names:		
Name:			Surname:	Surname:		
Addres	s:		DOB:	Address:		
			Address:			
Fax Nu	ımbe	r:		Fax Number:		
Vedo	lizuı	mab - continued				
Applic	ation	lication — Crohn's disease - children ns from any relevant practitioner. Approv ites(tick boxes where appropriate)				
	Paediatric patient has active Crohn's disease Patient has had an initial approval for prior biologic therapy and has experienced intolerable side effects or insufficient benefit to meet renewal criteria (unless contraindicated) Patient has a Paediatric Crohn's Disease Activity Index (PCDAI) score of greater than or equal to 30 or					
		Patient has extensive small intestine disease				
	and		perienced an inadequate response to (including lack of initial response and/or loss of initial response) immunomodulators and corticosteroids			
		Patient has experienced into	olerable side effects from immunomodulators and cor-	ticosteroids		
			ticosteroids are contraindicated			
Note:	Indic	cation marked with * is an unapproved in	ndication.			
Curre Applic	nt ap ation	— Crohn's disease - children* proval Number (if known): s from any relevant practitioner. Approvites (tick boxes where appropriate)				
		or PCDAI score is 15 or less	y 10 points from when the patient was initiated on bio			
	and	Vedolizumab to administered at a	dose no greater than 300mg every 8 weeks			
Note:	India	cation marked with * is an unapproved in	ndication.			

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Reg No:		First Names:	First Names:		
Name:		Surname:	Surname:		
Address	Σ	DOB:	Address:		
		Address:			
Fax Nun	mber:		Fax Number:		
Vedoli	zumab - continued				
Applica Prereq	meet renewal criteria (unless or Patient has a SCCAI score is patient's PUCAI score is gre Patient has tried but experie from prior therapy with immu or Patient has experienced into	proval for prior biologic therapy and has experienced contraindicated) s greater than or equal to 4	tial response and/or loss of initial response)		
Note: I	Indication marked with * is an unapproved in	dication.			
Current Applica	ral — ulcerative colitis t approval Number (if known): ations from any relevant practitioner. Approv				
	or	ed by 2 points or more from the SCCAI score since in ed by 10 points or more from the PUCAI score since			
		e no greater than 300 mg intravenously every 8 week	s		
Note: I	Note: Indication marked with * is an unapproved indication.				