Enquiries to Ministry of Health 0800 855 066

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPL	ICAN	T (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:				
Reg No:			First Names:	First Names:				
Name	e:		Surname:	Surname:				
Address:			DOB:	Address:				
			Address:					
Fax Number:				Fax Number:				
Ved	olizur	nab						
Appl	Initial application — Crohn's disease - adults Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)							
	and	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 CDAI score of greater than or equal to 300, or HBI score of greater than or equal to 10 Patient has extensive small intestine disease affecting more than 50 cm of the small intestine Patient has evidence of short gut syndrome or would be at risk of short gut syndrome with further bowel resection Patient has an ileostomy or colostomy, and has intestinal inflammation Patient has tried but experienced an inadequate response to (including lack of initial response and/or loss of initial response) from prior therapy with immunomodulators and corticosteroids Patient has experienced intolerable side effects from immunomodulators and corticosteroids Immunomodulators and corticosteroids are contraindicated						
Curr Appl	ent appication	or therapy CDAI score is 150 or less, o	vals valid for 2 years. 100 points, or HBI score has reduced by 3 points, from					
	and [Vedolizumab to administered at a	dose no greater than 300 mg every 8 weeks					

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			Address:			
Fax Number:				Fax Number:		
Vedo	lizur	mab - continued				
Applic	ation	lication — Crohn's disease - children as from any relevant practitioner. Approvites(tick boxes where appropriate)	als valid for 6 months.			
	and	Paediatric patient has active Crohr	's disease			
		or meet renewal criteria (unless	oroval for prior biologic therapy and has experienced contraindicated) nn's Disease Activity Index (PCDAI) score of greater			
		or Patient has extensive small i	iall intestine disease			
	and					
		Patient has tried but experier from prior therapy with immu	Patient has tried but experienced an inadequate response to (including lack of initial response and/or loss of initial response) from prior therapy with immunomodulators and corticosteroids			
		Patient has experienced into	lerable side effects from immunomodulators and cort	ticosteroids		
			icosteroids are contraindicated			
Note:	Indic	cation marked with * is an unapproved in	dication.			
Curre Applic	nt ap	- 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 an adequate response to treatment, but CDAI score			
	and [dose no greater than 300mg every 8 weeks			
Note:	India	cation marked with * is an unapproved in	dication			

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Name:			Surname:	Surname:			
Address:			DOB:	Address:			
			Address:				
				Fax Number:			
Vedolizumab - continued Initial application — ulcerative colitis Applications from any relevant practitioner. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)							
	and	Patient has had an initial appropriate or Patient has a SCCAI score is	Patient has a SCCAI score is greater than or equal to 4				
	and	Patient has tried but experienced an inadequate response to (including lack of initial response and/or loss of initial response) from prior therapy with immunomodulators and corticosteroids Patient has experienced intolerable side effects from immunomodulators and corticosteroids Immunomodulators and corticosteroids are contraindicated					
Note:	Indic	cation marked with * is an unapproved in	dication.				
Renewal — ulcerative colitis Current approval Number (if known):							
		or	ed by 2 points or more from the SCCAI score since in ed by 10 points or more from the PUCAI score since				
	and		e no greater than 300 mg intravenously every 8 week	s			
Note:	ote: Indication marked with * is an unapproved indication.						