Enquiries to Ministry of Health 0800 855 066

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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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:Bendamustine hydrochloride		Fax Number:						
Initial application — CLL* Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 12 months. Prerequisites(tick boxes where appropriate)								
The patient has chronic lymphocyl	tic leukaemia requiring treatment							
Patient has ECOG performance st	tatus of 0-2							
Bendamustine is to be administered	ed at a maximum dose of 100 mg/m² on days 1 and 2	every 4 weeks for a maximum of 6 cycles						
Note: Indication marked with a * includes indications that are unapproved. 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma (SLL).								
Initial application — Indolent, Low-grade lymphomas Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 9 months. Prerequisites(tick boxes where appropriate) The patient has indolent low grade NHL requiring treatment								
The patient has ECOG performan	The patient has ECOG performance status of 0-2							
Patient is treatment no and		ation with rituaireah when CD20						
or	e administered for a maximum of 6 cycles (in combina	ation with rituximab when GD20+)						
	or has relapsed within 12 months of a rituximab con	taining combined chemo-immunotherapy						
	e administered in combination with obinutuzumab for	a maximum of 6 cycles						
and Bendamustine is to be	eceived prior bendamustine therapy e administered for a maximum of 6 cycles in relapsed	patients (in combination with rituximab when						
and Patient has had a ritu:	ximab treatment-free interval of 12 months or more							
Bendamustine is to be administered as monotherapy for a maximum of 6 cycles in rituximab refractory patients								

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					Address:				
Fax N	umbe	r:				Fax Number:			
Bendamustine hydrochloride - continued									
Curre Appli	ent ap	proval is only	Num from	Low-grade lymphomas ber (if known): a relevant specialist or any xes where appropriate)	relevant practitioner on the recommendation of a rele	vant specialist. Approvals valid for 9 months.			
		and	Patient is refractory to or has relapsed within 12 months of rituximab in combination with bendamustine Bendamustine is to be administered in combination with obinutuzumab for a maximum of 6 cycles						
	or	Patients have not received a bendamustine regimen within the last 12 months and Bendamustine is to be administered for a maximum of 6 cycles in relapsed patients (in combination w when CD20+)							
				Patient has had	a rituximab treatment-free interval of 12 months or m	nore			
			or	Bendamustine is to be	e administered as a monotherapy for a maximum of 6	cycles in rituximab refractory patients			
Note	: 'indo	olent, l	ow-gr	rade lymphomas' includes fo	llicular, mantle cell, marginal zone and lymphoplasma	acytic/ Waldenstrom's macroglobulinaemia.			
Initial application — Hodgkin's lymphoma* Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)									
	and	Patient has Hodgkin's lymphoma requiring treatment							
	Patient has a ECOG performance status of 0-2 and Patient has received one prior line of chemotherapy and								
	and	Patient's disease relapsed or was refractory following prior chemotherapy							
				amustine is to be administere n/m2 twice per cycle, for a ma	ed in combination with gemcitabine and vinorelbine (Baximum of four cycles	BeGeV) at a maximum dose of no greater than			
Note	lote: Indications marked with * are unapproved indications.								

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