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.

	PATIENT:
lame:	
/ard:	NHI:
endamustine hydro	ochloride
NITIATION – CLL* Prerequisites (tick boxes	where appropriate)
and Patient ha	nt has chronic lymphocytic leukaemia requiring treatment as ECOG performance status 0-2
	stine is to be administered at a maximum dose of 100 mg/m² on days 1 and 2 every 4 weeks for a maximum of 6 cycles
lote: Indication marked v SLL).	with a * includes indications that are unapproved. 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma
and	
and	Patient is treatment naive Bendamustine is to be administered for a maximum of 6 cycles (in combination with rituximab when CD20+)
or O	Patient is refractory to or has relapsed within 12 months of a rituximab containing combined chemo-immunotherapy regimen Bendamustine is to be administered in combination with obinutuzumab for a maximum of 6 cycles
or O and	The patient has not received prior bendamustine therapy Bendamustine is to be administered for a maximum of 6 cycles in relapsed patients (in combination with rituximab when CD20+)
and	Patient has had a rituximab treatment-free interval of 12 months or more

I confirm that the above details are correct:	
Signed:	Date:

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.

ESCRIBER	PATIENT:	
me:		
rd:	NHI:	
ndamustine hydrochloride -	continued	
ONTINUATION – Indolent, Low-grad -assessment required after 9 months erequisites (tick boxes where approp		
and	ry to or has relapsed within 12 months of rituximab in combination with bendamustine to be administered in combination with obinutuzumab for a maximum of 6 cycles	
or O Patients have not and	received a bendamustine regimen within the last 12 months	
rituxim and	amustine is to be administered for a maximum of 6 cycles in relapsed patients (in combination with nab when CD20+) It has had a rituximab treatment-free interval of 12 months or more	
or Bendamusti	ine is to be administered as a monotherapy for a maximum of 6 cycles in rituximab refractory patients	
te: 'indolent, low-grade lymphomas' i	includes follicular, mantle cell, marginal zone and lymphoplasmacytic/ Waldenström's macroglobulinaemia.	
TIATION – Hodgkin's lymphoma* -assessment required after 6 months erequisites (tick boxes where approp		
O Patient has Hodgkin's lyi	mphoma requiring treatment	
O Patient has a ECOG per	Patient has a ECOG performance status of 0-2	
	e prior line of chemotherapy	
O Patient's disease relapse	ed or was refractory following prior chemotherapy	
O Bendamustine is to be a	administered in combination with gemcitabine and vinorelbine (BeGeV) at a maximum dose of no greater than le, for a maximum of four cycles	
te: Indications marked with * are una		

0:	D - 1
Signed.	Date.
Oldi Ica	