Enquiries to Ministry of Health 0800 855 066

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

Page 1 Form SA2443 June 2025

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:				Fax Number:		
Atezo	olizuma	b				
Appli	cations or	(tick boxes where appropriate) Patient has locally advanced or me	relevant practitioner on the recommendation of a me			
	and		ed treatment with an immune checkpoint inhibitor for stology there is documentation confirming that the diss not possible to ascertain			
	and Patient has an ECOG 0-2 and					
Patient has documented disease progression following treatment with at least two cycles		s of platinum-based chemotherapy				
Atezolizumab is to be used as monotherapy at a dose of 1200 mg every three weeks (or equivalent) for a				r equivalent) for a maximum of 16 weeks		
		Baseline measurement of overall to	umour burden is documented clinically and radiologic	ally		
Renewal — non-small cell lung cancer second line monotherapy Current approval Number (if known):						
	or		complete response to treatment			
	or	Patient's disease has had a partial response to treatment				
		Patient has stable disease				
	and	Response to treatment in target lesions has been determined by comparable radiologic assessment following the most recent treatment period				
	and	No evidence of disease progression	n			
	and	The treatment remains clinically ap	propriate and patient is benefitting from treatment			
	and	Atezolizumab to be used at a maxi	mum dose of 1200 mg every three weeks (or equival	ent)		
		Treatment with atezolizumab to cease 3 weeks)	ase after a total duration of 24 months from commen	cement (or equivalent of 35 cycles dosed every		

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

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	Address:	
Fax Number:		Fax Number:
Patient has locally advanced and Patient has preserved liver function and Transarterial chemoembolisa and Patient has not received or Patient received funde or Patient has expense.	th atezolizumab and met all remaining criteria prior to or metastatic, unresectable hepatocellular carcinomunction (Child-Pugh A) ation (TACE) is unsuitable and prior systemic therapy for the treatment of hepatocel deprior systemic therapy for the treatment of hepatocel deprior is unsuitable and prior systemic therapy for the treatment of hepatocel deprior is systemic therapy for the treatment of hepatocel deprivation before 1 March 2025 arienced treatment-limiting toxicity from treatment with gression since initiation of lenvatinib	a cellular carcinoma
Renewal — unresectable hepatocellular carcino Current approval Number (if known):	als valid for 6 months.	

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