Enquiries to Ministry of Health 0800 855 066

## APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

Page 1 Form SA2270 July 2025

		04., 2020
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:
Sirolimus (Rapamune)		
Prerequisites(tick box where appropriate)  The drug is to be used for rescue thera	ovals valid without further renewal unless notified.  py for an organ transplant recipient o calcineurin inhibitor treatment as defined by refracto	ry rejection; or intolerant to calcineurin inhibitor
• GFR< 30 ml/min; or		
Rapidly progressive transplant vasculopathy;	or	
Rapidly progressive obstructive bronchiolitis;	or	
HUS or TTP; or		
Leukoencepthalopathy; or		
Significant malignant disease		
Initial application — severe non-malignant lyr Applications from any relevant practitioner. Appr Prerequisites(tick boxes where appropriate)		
Patient has severe non-malignan	t lymphovascular malformation*	
or	quately controlled by sclerotherapy and surgery ead/extensive and sclerotherapy and surgery are not	considered clinically appropriate
or Sirolimus is to be used to r	educe malformation prior to consideration of surgery	
and	cialist lymphovascular malformation multi-disciplinary	eam

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

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Sirolimus (Rapamune) - continued						
Renewal — severe non-malignant lymphovascu Current approval Number (if known): Applications from any relevant practitioner. Approv Prerequisites(tick boxes where appropriate)						
Patient's disease has had either a complete response or a partial response to treatment, or patient has stable disease according to RECIST version 1.1 (see Note)  Patient's disease has stabilised or responded clinically and disease response to treatment has been clearly documents in patient notes  and  No evidence of progressive disease and  The treatment remains clinically appropriate and the patient is benefitting from the treatment  Note: Baseline assessment and disease responses to be assessed according to the Response Evaluation Criteria in Solid Tumours (RECIST) version 1.1 (Eisenhauer et al. Eur J Cancer 2009;45:228-47) Indications marked with * are unapproved indications						
Initial application — renal angiomyolipoma(s) associated with tuberous sclerosis complex* Applications only from a nephrologist or urologist. Approvals valid for 6 months.  Prerequisites(tick boxes where appropriate)						
and Patient has tuberous sclerosis com  Evidence of renal angiomyolipoma	nplex* (s) measuring 3 cm or greater and that have shown i	nterval growth				
Renewal — renal angiomyolipoma(s) associate Current approval Number (if known): Applications from any relevant practitioner. Approv Prerequisites(tick boxes where appropriate)						
and Demonstrated stabilisation or impre	giomyolipoma reduction or stability by magnetic resor ovement in renal function ngiomyolipoma haemorrhage or significant adverse e					
The treatment remains appropriate  Note: Indications marked with * are unapproved in	and the patient is benefitting from treatment					

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Address:			DOB:	Address:	
				Address:	
Fax Numbe	er:		<b>1e)</b> - continued		Fax Number:
Initial app	olications only	on — ref y from a i ck boxes	ractory seizures assoc neurologist. Approvals va where appropriate)	iated with tuberous sclerosis complex* alid for 6 months.  round of documented tuberous sclerosis complex	
	or	Vigabatrin has been trialled and has not adequately controlled seizures  and  Seizures are not adequately controlled by, or the patient has experienced unacceptable side effects from, optimal treatment with at least two of the following: sodium valproate, topiramate, levetiracetam, carbamazepine, lamotrigine, phenytoin sodium, and lacosamide (see Note)			
		and	treatment with at leas	dicated quately controlled by, or the patient has experienced ut three of the following: sodium valproate, topiramate d lacosamide (see Note)	
and		Patient ha	have a significant impactass been assessed and so DR inhibitor treatment pri	urgery is considered inappropriate for this patient, or t	he patient has been assessed and would benefit
			ing age potential are not m valproate.	required to trial phenytoin sodium, sodium valproate,	or topiramate. Those who can father children are
Current ap Application Prerequis	oprova ns only <b>sites</b> (ti	I Number y from a r ck box w	r (if known): neurologist. Approvals va here appropriate)		oizura fraguanov) ar covarity and/ar nationt availth.
(	of life o	compared		improvement in seizure rate (e.g. 50% reduction in sarting sirolimus treatment ndications	eizure frequericy) or severity and/or patient quality

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