APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

Page 1 Form SA2254 November 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:			
Bosentan					
cardiologist or rheumatologist. Approvals valid for or Prerequisites (tick boxes where appropriate) Patient has pulmonary arterial hyperand PAH is in Group 1, 4 or 5 of the Whand					
and A mean pulmonary art and A pulmonary capillary and Pulmonary vascular re and PAH has been d defined in the 20 or Patient has not e risk stratification	ed by right heart catheterisation ery pressure (PAPm) greater than 20 mmHg (unless wedge pressure (PCWP) less than or equal to 15 mm sistance greater than 2 Wood Units or greater than 1 temonstrated to be non-responsive in vasoreactivity a 222 ECS/ERS Guidelines for PAH (see note below for experienced an acceptable response to calcium anta tool**	nHg 60 International Units (dyn s cm ⁻⁵) assessment using iloprost or nitric oxide, as r link to these guidelines) † gonist treatment, according to a validated			
or disorders including severe ch	econdary to congenital heart disease or PAH due to in pronic neonatal lung disease entricle congenital heart disease and elevated pulmonhe minimising of pulmonary/venous filling pressures				
Bosentan is to be used as PA	Bosentan is to be used as PAH monotherapy				
Patient has experienced intolerable side effects on sildenafil or					
Patient has an absolut	e contraindication to sildenafil				
Patient is a child with in	diopathic PAH or PAH secondary to congenital heart	disease			

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 Numbe	er:				Fax Number:
Bosentai	n - co	ontinue	ed		
Application cardiologis	ns onl	y from heuma ick box Patien	atologist. Approvals valid for xes where appropriate) It has pulmonary arterial hype in Group 1, 4 or 5 of the West in New York Heart Association. PAH has been confirm A mean pulmonary and A pulmonary capillary. Pulmonary vascular results.		onal class II, III or IV s peri Fontan repair) nmHg 160 International Units (dyn s cm ⁻⁵)
	or or	_	Patient has not risk stratificatio Patient has PAI Patient is a child with PAH s disorders including severe of	H other than idiopathic / heritable or drug-associated secondary to congenital heart disease or PAH due to chronic neonatal lung disease	agonist treatment, according to a validated type idiopathic, congenital or developmental lung
and				ventricle congenital heart disease and elevated pulm the minimising of pulmonary/venous filling pressures	
and	and Bosentan is to be used as part of PAH dual therapy and				
	or			notherapy (sildenafil) for at least three months and harding to a validated risk stratification tool**	as experienced an inadequate therapeutic
			Patient is presenting in NYF initial dual therapy	HA/WHO functional class III or IV, and in the opinion of	of the treating clinician would likely benefit from

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Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Bosentan - continued		
Patient has pulme and PAH is in New You and PAH and A me	ory specialist, cardiologist, rheumatologist or any relevant practiti pprovals valid for 6 months.	functional class II, III or IV
or Patient is a disorders in Or Patient has	Imonary capillary wedge pressure (PCWP) less than or equal to nonary vascular resistance greater than 2 Wood Units or greater PAH has been demonstrated to be non-responsive in vasoread defined in the 2022 ECS/ERS Guidelines for PAH (see note be Patient has not experienced an acceptable response to calciur risk stratification tool** Patient has PAH other than idiopathic / heritable or drug-associated with PAH secondary to congenital heart disease or PAH decluding severe chronic neonatal lung disease	ctivity assessment using iloprost or nitric oxide, as elow for link to these guidelines) † m antagonist treatment, according to a validated ciated type due to idiopathic, congenital or developmental lung
and Bosentan is and Patie	ent is on the lung transplant list ent is presenting in NYHA/WHO functional class IV Patient has tried PAH dual therapy for at least three months ar treatment according to a validated risk stratification tool** Patient does not have major life-threatening comorbidities and	nd has not experienced an acceptable response to
	scenario	a upic arcrapy is not being used in a palliative

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Name:	Surname:	Surname:			
Address:	DOB:	Address:			
	Address:				
Fax Number:		Fax Number:			
Bosentan - continued					
Renewal					
Current approval Number (if known):					
Applications only from a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation of a respiratory specialist, cardiologist or rheumatologist. Approvals valid for 2 years. Prerequisites(tick box where appropriate)					
Patient is continuing to derive benefit from bosentan treatment according to a validated PAH risk stratification tool**					

Note: † The European Respiratory Journal Guidelines can be found here: 2022 ECS/ERS Guidelines for the diagnosis and treatment of pulmonary

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hypertension PAH

** the requirement to use a validated risk stratification tool to determine insufficient response applies to adults. Determining insufficient response in children does not require use of a validated PAH risk stratification tool, where currently no such validated tools exist for PAH risk stratification in children.