RESCRIBER			PATIENT:
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nbrise	ntan		
mbriser NITIATION Re-assess Prerequis	ntan N - PA sment i sites (ti Prescri a respi Hospita O F	AH mon required ick boxe bed by, ratory spal.  Patient h PAH is ir  and and and and PAH is represented the representation of the rep	NHI:
and	or		atient has palliated single ventricle congenital heart disease and elevated pulmonary pressures or a major complication of the intan circulation requiring the minimising of pulmonary/venous filling pressures
	and	) An	mbrisentan is to be used as PAH monotherapy
		or _	Patient has experienced intolerable side effects with both sildenafil and bosentan
			Patient has an absolute contraindication to sildenafil and an absolute or relative contraindication to bosentan (e.g. due to current use of a combined oral contraceptive or liver disease)
		or	

CRIBER	PATIENT:
:	Name:
	NHI:
risentan	- continued
ssessment	AH dual therapy required after 6 months ick boxes where appropriate)
	bed by, or recommended by a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation ratory specialist, cardiologist or rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health Nal.
O F	Patient has pulmonary arterial hypertension (PAH)
	PAH is in Group 1, 4 or 5 of the WHO (Venice 2003) clinical classifications
	PAH is in New York Heart Association/World Health Organization (NYHA/WHO) functional class II, III or IV
	PAH has been confirmed by right heart catheterisation
	A mean pulmonary artery pressure (PAPm) greater than 20 mmHg (unless peri Fontan repair)
	A pulmonary capillary wedge pressure (PCWP) less than or equal to 15 mmHg
	Pulmonary vascular resistance greater than 2 Wood Units or greater than 160 International Units (dyn s cm <sup>-5</sup> )
	PAH has been demonstrated to be non-responsive in vasoreactivity assessment using iloprost or nitric oxide, as defined in the 2022 ECS/ERS Guidelines for PAH (see note below for link to these guidelines) †
	Patient has not experienced an acceptable response to calcium antagonist treatment, according to a validated risk stratification tool**
	O Patient has PAH other than idiopathic / heritable or drug-associated type
or	Patient is a child with PAH secondary to congenital heart disease or PAH due to idiopathic, congenital or developmental lung disorders including chronic neonatal lung disease
	<ul> <li>Patient has palliated single ventricle congenital heart disease and elevated pulmonary pressures or a major complication of the Fontan circulation requiring the minimising of pulmonary/venous filling pressures</li> </ul>
and	O Ambrisentan is to be used as PAH dual therapy
and	Patient has tried a PAH monotherapy (sildenafil or bosentan) for at least three months and has not experienced an acceptable response to treatment according to a validated risk stratification tool**
	O Patient has tried PAH dual therapy including bosentan and has experienced intolerable side effects on bosentan
and	
	Patient is presenting in NYHA/WHO functional class III or IV, and in the opinion of the treating clinician would benefit from initial dual therapy
	Patient has an absolute or relative contraindication to bosentan (eg due to current use of a combined oral contraceptive

I confirm that the above details are correct:

Signed: Date:

SCRIBER		PATIENT:				
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brisentan	- continued					
assessment r	AH triple thera required after 6 ck boxes wher	5 months				
	ratory specialis	ommended by a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation of st, cardiologist or rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ				
and	•	monary arterial hypertension (PAH) o 1, 4 or 5 of the WHO (Venice 2003) clinical classifications				
and		York Heart Association/World Health Organization (NYHA/WHO) functional class II, III or IV				
	and	H has been confirmed by right heart catheterisation nean pulmonary artery pressure (PAPm) greater than 20 mmHg (unless peri Fontan repair)				
	and A p	oulmonary capillary wedge pressure (PCWP) less than or equal to 15 mmHg				
	and Puli	monary vascular resistance greater than 2 Wood Units or greater than 160 International Units (dyn s cm <sup>-5</sup> )				
	or	PAH has been demonstrated to be non-responsive in vasoreactivity assessment using iloprost or nitric oxide, as defined in the 2022 ECS/ERS Guidelines for PAH (see note below for link to these guidelines) †				
	or	Patient has not experienced an acceptable response to calcium antagonist treatment, according to a validated risk stratification tool**				
		Patient has PAH other than idiopathic / heritable or drug-associated type				
or (		a child with PAH secondary to congenital heart disease or PAH due to idiopathic, congenital or developmental lung including chronic neonatal lung disease				
		as palliated single ventricle congenital heart disease and elevated pulmonary pressures or a major complication of the rculation requiring the minimising of pulmonary/venous filling pressures				
and (and	O Ambrisen	ntan is to be used as PAH triple therapy				
	O Patient is on the lung transplant list  or					
	and	Patient is presenting in NYHA/WHO functional class IV				
		Patient has an absolute or relative contraindication to bosentan (e.g. due to current use of a combined oral contraceptive or liver disease)				
	or and	Patient has tried PAH dual therapy for at least three months and remains in an unacceptable risk category according to a validated risk stratification tool**				
		Patient does not have major life-threatening comorbidities and triple therapy is not being used in a palliative scenario				

I confirm that the above details are correct:

Signed: Date:

PRESCR	IBER	PATIENT:						
Name:		Name:						
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Ambrisentan - continued								
CONTINUATION Re-assessment required after 2 years Prerequisites (tick box where appropriate)								
and	Prescribed by, or recommended by a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation of a respiratory specialist, cardiologist or rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital.  The patient is continuing to derive benefit from ambrisentan 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 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.

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