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.

PRESCRIBER	PATIENT:
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oprost	
Re-assessmen Prerequisites Preso	PAH monotherapy t required after 6 months (tick boxes where appropriate) pribed by, or recommended by a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation of biratory specialist, cardiologist or rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ ital.
and and and and and and	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 and A mean pulmonary artery pressure (PAPm) greater than 20 mmHg (unless peri Fontan repair) and A pulmonary capillary wedge pressure (PCWP) less than or equal to 15 mmHg and
	A pulmonary vascular resistance greater than 2 Wood Units or greater than 160 International Units (dyn s cm ⁻⁵) A pulmonary vascular resistance greater than 2 Wood Units or greater than 160 International Units (dyn s cm ⁻⁵) 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** 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 severe chronic neonatal lung disease 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
and	O Iloprost is to be used as PAH monotherapy O Patient has experienced intolerable side effects on sildenafil and both the funded endothelin receptor antagonists (i.e. both bosentan and ambrisentan)

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

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ssessment equisites (t require (tick bo ribed b piratory	al therapy ed after 6 months xes where appropriate) y, or recommended by a respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation specialist, cardiologist or rheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health N
and		t has pulmonary arterial hypertension (PAH)
and		in Group 1, 4 or 5 of the WHO (Venice 2003) clinical classifications in New York Heart Association/World Health Organization (NYHA/WHO) functional class II, III or IV
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or	0	Patient is a child with PAH secondary to congenital heart disease or PAH due to idiopathic, congenital or developmental lung disorders including severe chronic neonatal lung disease 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
and		loprost is to be used as PAH dual therapy with either sildenafil or an endothelin receptor antagonist
	or	Patient has an absolute contraindication to or has experienced intolerable side effects on sildenafil Patient has an absolute or relative contraindication to or experienced intolerable side effects with a funded endothelin receptor antagonist
and	t	O Patient has tried a PAH monotherapy for at least three months and remains in an unacceptable risk category according

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	respiratory specialist, cardiologist, rheumatologist or any relevant practitioner on the recommendation rrheumatologist, or in accordance with a protocol or guideline that has been endorsed by the Health N
Patient has pulmonary arterial hand PAH is in Group 1, 4 or 5 of the	nypertension (PAH) WHO (Venice 2003) clinical classifications
and	ciation/World Health Organization (NYHA/WHO) functional class II, III or IV
and A mean pulmonary and A pulmonary capilla and A pulmonary vascul and PAH has been defined in the or Patient has no risk stratificati or Patient has	AH other than idiopathic / heritable or drug-associated type
and O Patient has palliated single Fontan circulation requiring and O Illoprost is to be used as Formation and O Patient is on the lunger or O Patient is presenting or O Patient has trivial and O Pa	
and	not have major life-threatening comorbidities and triple therapy is not being used in a palliative

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PRESCRIBER	PATIENT:
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	gist, rheumatologist or any relevant practitioner on the recommendation of nce with a protocol or guideline that has been endorsed by the Health NZ ing to a validated PAH risk stratification tool
Note: † The European Respiratory Journal Guidelines can be found here: diagnosis and treatment of pulmonary hypertension PAH ** the requirement to use a validated risk stratification tool to determine ins Determining insufficient response in children does not require use of a validate currently no such validated tools exist for PAH risk stratification in children.	ufficient response applies to adults.

I confirm that the above details are correct:

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