

medicare



Paroxysmal nocturnal haemoglobinuria – eculizumab or ravulizumab – initial grandfather authority application

Online PBS Authorities

You do not need to complete this form if you use the Online PBS Authorities system.

For more information and how to access the **Online PBS Authorities** system, go to **servicesaustralia.gov.au/hppbsauthorities**

When to use this form

Use this form to apply for **initial grandfather** PBS-subsidised eculizumab or ravulizumab for patients with paroxysmal nocturnal haemoglobinuria (PNH) who had, prior to **1 March 2022**:

- received non-PBS-subsidised treatment with eculizumab or ravulizumab maintenance phase
- received eculizumab through the Australian Government's Life Saving Drugs Program (LSDP) and are continuing on eculizumab.

Important information

Initial grandfather applications to start PBS-subsidised treatment can be made using the **Online PBS Authorities** system or in writing and must include sufficient information to determine the patient's eligibility according to the PBS criteria.

Under no circumstances will phone approvals be granted for PNH **initial grandfather** authority applications.

Complement 5 (C5) inhibitors are defined as eculizumab or ravulizumab.

The information in this form is correct at the time of publishing and may be subject to change.

Continuing treatment

This form is ONLY for initial grandfather treatment.

For **continuing** PBS-subsidised treatment, the patient must qualify under the **first continuing** or **subsequent continuing** treatment criteria.

Section 100 arrangements for eculizumab and ravulizumab

These items are available to a patient who is attending:

- an approved private hospital, or
- a public hospital

and is a:

- · day admitted patient
- non-admitted patient, or
- patient on discharge.

These items are not available as a PBS benefit for in-patients of a public hospital.

The hospital name and provider number must be included in this authority form.

For more information

Go to servicesaustralia.gov.au/healthprofessionals

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You do not need to complete this form if you use the Online PBS Authorities system. Go to servicesaustralia.gov.au/hppbsauthorities Patient's details Medicare card number Department of Veterans' Affairs card number Mrs | Miss | Ms Family name First given name 3 Date of birth (DD MM YYYY) Patient's current weight Prescriber's details 5 Prescriber number Mrs Miss Ms __ Mr Other Family name First given name 7 Business phone number (including area code) Alternative phone number (including area code)

Ho	spital details					
8	Hospital name					
	This hospital is a:					
	public hospital					
	private hospital					
9	Hospital provider number					
Co	nditions and criteria					
	qualify for PBS authority approval, the following conditions ust be met.					
10	The patient is being treated by a:					
	haematologist					
	non-specialist medical physician who has consulted a haematologist					
11	This application is for:					
	eculizumab Go to 12					
	ravulizumab Go to 14					
12	The patient:					
	has received non-PBS-subsidised treatment with eculizumab for this condition prior to 1 March 2022					
	or					
	has previously received eculizumab for the treatment of this condition funded under the Australian Government's LSDP.					
13	Has the patient experienced clinical improvement or a stabilisation of the condition as a result of treatment with this					
	drug?					
	Yes Go to 16					
	No .					
14	Has the patient received non-PBS-subsidised treatment with ravulizumab for this condition prior to 1 March 2022?					
	Yes					
	No 🗌					



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IJ	Has the patient demonstrated clinical improvement or	19	Provide details of th	ie following r	nonitorinç	g requir	ements
	stabilisation of the condition with details kept with the patient's record?		Test	Result	Date o	f test ([DD MM YYYY)
	Yes		Haemoglobin (g/L)				
	No 🗌		Platelets (x10 ⁹ /L)				
16	Is this treatment in combination with another C5 inhibitor or pegcetacoplan?		White Cell Count (x10 ⁹ /L)				
	Yes No		Reticulocytes (x10 ⁹ /L)				
17	Prior to commencing treatment with this drug, the patient had:		Neutrophils (x10 ⁹ /L)				
	a diagnosis of PNH established by flow cytometry		Granulocyte clone size (%)				
	a PNH granulocyte clone size ≥ 10%		LDH				
	and		ULN for LDH as				
	a raised lactate dehydrogenase (LDH) value at least 1.5 times the upper limit of normal (ULN).		quoted by the reporting laboratory				
18	Prior to commencing treatment with this drug, the patient had:		LDH : ULN ratio (in figures, rounded				
	experienced a thrombotic/embolic event which required anticoagulant therapy		to one decimal place & must be				
	or		at least 1.5)				
	been transfused with at least 4 units of red blood cells in the previous 12 months	Che	ecklist				
	or	20	The releva	nt attachmer	nts need t	o be pro	ovided with
	debilitating shortness of breath/chest pain resulting in limitation of normal activity (New York Heart Association Class III) and/or established diagnosis of pulmonary arterial hypertension, where causes other than PNH had been	this form. Details of the proposed prescription(s).					
	excluded	Priv	acy notice				
	or	21	Personal informatio	n is protecte	d hy law ((includir	na the
	□ a history of renal insufficiency, demonstrated by an eGFR ≤ 60 mL/min/1.73m ² , where causes other than PNH had been excluded		Privacy Act 1988) a purposes of assessi	nd is collecteing and proce	ed by Servessing this	vices Au s autho	ustralia for the rity application.
	or		Personal informatio	•	•		,
	recurrent episodes of severe pain requiring hospitalisation and/or narcotic analgesia where causes other than PNH had been excluded		given to other partion where it is required purpose of research	or authorise	d by law ((includir	ng for the
	or		More information at				
	chronic/recurrent anaemia, where causes other than haemolysis had been excluded, together with multiple haemoglobin measurements:		manages personal i be found at service	,			
	not exceeding 70 g/L in the absence of anaemia symptoms						
	or						
	not exceeding 100 g/L in addition to having anaemia symptoms.						

Prescriber's declaration

You do not need to **sign** the declaration if you complete this form using Adobe Acrobat Reader and return this form through Health Professional Online Services (HPOS) at

servicesaustralia.gov.au/hpos

22 I declare that:

- I am aware that this patient must meet the criteria listed in the current Schedule of Pharmaceutical Benefits to be eligible for this medicine.
- I have informed the patient that their personal information (including health information) will be disclosed to Services Australia for the purposes of assessing and processing this authority application.
- I have provided details of the proposed prescription(s) and the relevant attachments as specified in the Pharmaceutical Benefits Scheme restriction.
- the information I have provided in this form is complete and correct.

I understand that:

• giving false or misleading information is a serious offence.					
I have read, understood and agree to the above.					
Date (DD MM YYYY) (you must date this declaration)					
Prescriber's signature (only required if returning by post)					

Returning this form

Return this form, details of the proposed prescription(s) and any relevant attachments:

 online (no signature required), upload through HPOS at servicesaustralia.gov.au/hpos

nr

 by post (signature required) to Services Australia

> Complex Drugs Programs Reply Paid 9826

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