

medicare



Severe asthma – adolescent and adult – initial or recommencement after 12 months break 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** or **recommencing** PBS-subsidised biological medicines for patients 12 years or over with uncontrolled severe asthma.

Important information

Initial or **recommencement** 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.

Applications for **balance of supply** can be made in real time using the **Online PBS Authorities** system or by phone. Call 1800 700 270 Monday to Friday, 8 am and 5 pm, local time.

Under no circumstances will phone approvals be granted for uncontrolled severe asthma **initial** or **recommencement** authority applications.

Where the term 'biological medicine' appears, it refers to benralizumab, dupilumab, mepolizumab and omalizumab.

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 or recommencement of treatment.

Following the completion of an **initial** treatment course with a specific biological medicine, a patient may qualify to receive up to 24 weeks of **continuing** treatment with that biological medicine providing they have demonstrated an adequate response to treatment.

Applications for **continuing** treatment with benralizumab, dupilumab or mepolizumab 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.

Applications for **continuing** treatment with omalizumab can be made in real time using the **Online PBS Authorities** system or by phone. Call 1800 700 270 Monday to Friday, 8 am to 5 pm, local time.

Section 100 arrangements for benralizumab, dupilumab, mepolizumab and omalizumab

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.

Treatment specifics

The assessment of the patient's response to the course of treatment must be conducted within the time frame specified in the restriction. Where a demonstration of response is not conducted within the required time frame, the patient will be deemed to have failed treatment with that particular PBS-subsidised biological medicine.

A patient who has experienced a serious adverse reaction of a severity necessitating permanent treatment withdrawal is not considered to have failed treatment with that particular PBS-subsidised biological medicine.

The patient must not receive more than 32 weeks of treatment under this restriction.

For more information

Go to servicesaustralia.gov.au/healthprofessionals

PB075.2502



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Hospital details Online PBS Authorities You do not need to complete this form if you use the Hospital name Online PBS Authorities system. Go to servicesaustralia.gov.au/hppbsauthorities This hospital is a: __ public hospital Patient's details private hospital Medicare card number Hospital provider number Department of Veterans' Affairs card number **Conditions and criteria** To qualify for PBS authority approval, the following conditions must be met. Dr Mr Mrs Miss Ms The patient, 12 years or over, is being treated by a medical Family name practitioner who is: a respiratory physician First given name a clinical immunologist an allergist 3 Date of birth (DD MM YYYY) a general physician experienced in the management of patients with severe asthma. **10** The patient has been: under the care of the same physician for at least Prescriber's details 6 months Prescriber number or diagnosed by a multidisciplinary severe asthma clinic team. **11** The patient has: Mr Mrs Miss not received PBS-subsidised treatment with a biological Family name medicine for severe asthma had a break in treatment of at least 12 months from the First given name most recently approved PBS-subsidised biological medicine for severe asthma. Business phone number (including area code) 12 Will this treatment be used in combination with and within 4 weeks of another PBS-subsidised biological medicine for

13 Has the patient had asthma for at least 1 year?

MCA0PB075 2502

Alternative phone number (including area code)

severe asthma?

Yes No

Yes No

	he patient has a diagnosis of asthma, confirmed and	17 Provide details of contraindications and/or intolerances of a
a	ocumented in the patient's medical records by either:	severity necessitating permanent treatment withdrawal to standard therapy according to the relevant
	a respiratory physician a clinical immunologist	TGA-approved Product Information.
	a a allergist	For details of the toxicity criteria, go to
	a general physician experienced in the management of	servicesaustralia.gov.au/healthprofessionals
_	patients with severe asthma	Inhaled corticosteroid
	Go to 15	
0	¬	
	at least 2 physicians experienced in the management of patients with severe asthma.	
	Go to 16	Inhaled long acting beta-2 agonist therapy
15 ⊤	he patient's diagnosis was defined at the time by at least one	
	f the following standard clinical features:	
	forced expiratory volume (FEV1) reversibility ≥ 12% and	
	≥ 200 mL at baseline within 30 minutes after administration of salbutamol (200 to 400 mcg)	Oral corticosteroids (for dupilumab 300 mg applications only)
	airway hyperresponsiveness with FEV1 decline > 20%	
	during a direct bronchial provocation test	
	airway hyperresponsiveness with FEV1 decline > 15%	
	during an indirect bronchial provocation test	▶ Go to 19
L	peak expiratory flow (PEF) variability > 15% between the 2 highest and 2 lowest PEF rates during 14 days.	18 The patient has failed to achieve adequate control with
16 ⊤	he patient has:	optimised asthma therapy in the past 12 months, despite formal assessment of and adherence to correct inhaler technique,
	received optimised asthma therapy including:	which has been documented in the patient's medical records
	adherence to high dose inhaled corticosteroid (ICS) for	and demonstrated by:
	at least 12 months	at least one admission to hospital for a severe asthma exacerbation while receiving optimised asthma therapy
	From (DD MM YYYY)	Date of exacerbation (DD MM YYYY)
	To (DD MM YYYY)	or
	adherence to long acting beta-2 agonist (LABA)	at least one severe asthma exacerbation requiring
	therapy for at least 12 months	documented use of systemic corticosteroids prescribed or
	From (DD MM YYYY)	supervised by a physician with either:
		OCS initiated or increased for at least 3 days Date of exacerbation (DD MM YYYY)
	To (DD MM YYYY) and (for dupilumab 300 mg applications only)	Date of exacerbation (DD WIWI 1111)
	regular maintenance oral corticosteroids (OCS) in the	
	last 6 months with a stable daily OCS dose of 5 to 35	or parenteral corticosteroids
	mg/day of prednisolone or equivalent over the 4 weeks prior to treatment initiation	Date of exacerbation (DD MM YYYY)
	Go to 18	
0		19 Does the patient have a baseline Asthma Control Questionnaire
	contraindications and/or intolerances to prior optimised	(ACQ-5) score of ≥ 2.0 (no more than one month old)?
	asthma therapy	Yes
	Go to 17	No 🗆
		20 Provide baseline details
		ACQ-5 Score
		Date (DD MM YYYY)

21	This application is for:	Privacy notice
	Benralizumab Go to 23	27 December information is protected by law (including the
	Dupilumab Go to 22	27 Personal information is protected by law (including the <i>Privacy Act 1988</i>) and is collected by Services Australia for the
	Mepolizumab Go to 23	purposes of assessing and processing this authority application.
	Omalizumab Go to 24	Personal information may be used by Services Australia, or
22	Which qualifying blood test results will be provided with this authority application?	given to other parties where the individual has agreed to this, or where it is required or authorised by law (including for the purpose of research or conducting investigations).
	☐ Blood eosinophil count	More information about the way in which Services Australia
	☐ IgE level	manages personal information, including our privacy policy, can
23	In the last 12 months, the patient has had:	be found at servicesaustralia.gov.au/privacypolicy
	a baseline blood eosinophil count ≥ 150 cells/microlitre while receiving treatment with OCS	Prescriber's declaration
	Blood eosinophil count cells per microlitre	You do not need to sign the declaration if you complete this form
	Date (DD MM YYYY)	using Adobe Acrobat Reader and return this form through Health Professional Online Services (HPOS) at
	Go to 26	servicesaustralia.gov.au/hpos
	or (not applicable to dupilumab 300 mg applications)	28 I declare that:
	a baseline blood eosinophil count ≥ 300 cells/microlitre	I am aware that this patient must meet the criteria listed in
	Blood eosinophil count cells per microlitre	the current Schedule of Pharmaceutical Benefits to be eligible for this medicine.
04	Date (DD MM YYYY) Go to 26	 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.
2 4	In the last 12 months, the patient has had:	 I have provided details of the proposed prescription(s) and
	total serum human immunoglobulin E (lgE) ≥ 30 lU/mL with past or current evidence of atopy, documented by skin prick testing	the relevant attachments as specified in the Pharmaceutical Benefits Scheme restriction.
	or	the information I have provided in this form is complete and
	total serum human $IgE \ge 30 IU/mL$ with past or current	correct. I understand that:
	evidence of atopy, documented by an in vitro measure of specific IgE	 giving false or misleading information is a serious offence.
25		
23	Provide the patient's total serum human IgE (no more than 12 months old)	I have read, understood and agree to the above.
	IgE result	Date (DD MM YYYY) (you must date this declaration)
	Date (DD MM YYYY)	Prescriber's signature (only required if returning by post)
Ch	ecklist	
26	' '	
	this form.	Returning this form
	Details of the proposed prescription(s).	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
		or
		by post (signature required) to
		Services Australia Complex Drugs Programs
		Reply Paid 9826
		HOBART TAS 7001