





Severe asthma – adolescent and adult – change authority application

When to use this form	Use this form to apply for changing PBS-subsidised biological medicines for patients 12 years or over with uncontrolled severe asthma.				
Important information	Authority applications must be 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 to 5 pm, local time.				
	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 changing treatment.				
	Following the completion of a change of 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 must be 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	These items are available to a patient who is attending:				
for benralizumab, dupilumab, mepolizumab	an approved private hospital, or				
and omalizumab	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				



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Pa	tient's details	Conditions and criteria	
1	Medicare card number	To qualify for PBS authority approval, the following conditions must be met.	
2	or Department of Veterans' Affairs card number Dr Mr Mrs Miss Ms	 9 The patient is being treated by a medical practitioner who is: a respiratory physician a clinical immunologist an allergist a general physician experienced in the management of 	:
	Family name	patients with severe asthma. 10 The patient has been:	ntha
	First given name	 under the care of the same physician for at least 6 mor or diagnosed by a multidisciplinary severe asthma clinic te 	
3	Date of birth (DD MM YYYY)	11 Has the patient received prior PBS-subsidised treatment with biological medicine for severe asthma in this treatment cycle No	
Pr	escriber's details	Yes Provide details	
4	Prescriber number	Prior biological medicine	
5	Dr Mr Mrs Miss Miss Other	From (DD MM YYYY)	
6	First given name Business phone number (including area code)	12 Has the patient failed, or ceased to respond to, PBS-subsidis treatment with this drug (the biological medicine this applica is for) for severe asthma during the current treatment cycle?	ation
	Alternative phone number (including area code)	 13 Will this treatment be used in combination with and within 4 weeks of another PBS-subsidised biological medicine for severe asthma? 	
Ho	spital details	No Yes	
7	Hospital name		
	This hospital is a: public hospital private hospital		
8	Hospital provider number		
		MCA0PB285 2410	

14 The	e patient is switching biological medicine treatment due to: failure to demonstrate or sustain response to prior biological medicine treatment for severe asthma	16 The patient has demonstrated a response to the most recent PBS-subsidised biological medicine treatment for severe asthma, assessed no more than 4 weeks after the last dose of
	Go to 15	biological medicine and evidenced by:
or		a reduction in the ACQ-5 score of at least 0.5 from baseline
	partial response to prior biological medicine treatment for severe asthma	Current ACQ-5 Score
	Go to 16	Date of current score (DD MM YYYY)
or		
	experiencing an adverse event due to prior biological	
	medicine treatment for severe asthma	or a reduction in the maintenance dose of oral corticosteroid
	Go to 15	(OCS) by at least 25% from baseline
or		Name of steroid
	other reason	
		Current dose
		mg/day
		and
		no deterioration in the ACQ-5 score from baseline
	Go to 15	Current ACQ-5 Score
15 The	e patient:	
	is submitting a new baseline Asthma Control Questionnaire	Date of current score (DD MM YYYY)
	(ACQ-5) score of:	
	and if applicable, is receiving maintenance oral	or
	corticosteroids (OCS) dose of:	an increase of up to 0.5 in the ACQ-5 score from baseline
	mg/day	Current ACQ-5 Score
	ing, duy	
	and	
	an assessment of response will be conducted around	Date of current score (DD MM YYYY)
	28 weeks after the first dose of this treatment	
or		17 This application is for:
	is using the previously submitted baseline ACQ-5 score of:	Benralizumab
		Dupilumab 200 mg Go to 20
	and	Dupilumab 300 mg
	future demonstrations of response will be assessed	Mepolizumab
	against the previously recorded baseline	Omalizumab Go to 22
	• Go to 17	
		18 Has the patient been receiving regular maintenance OCS in the last 6 months with a stable daily OCS dose of 5 to 35 mg/day of prednisolone or equivalent over the 4 weeks prior to treatment initiation?
		Yes
		Not applicable by due to contraindications or intolerances of
		a severity necessitating permanent treatment withdrawal
		19 If applicable, provide details of contraindications and/or intolerances to the regular maintenance OCS therapy.

	authority application? Blood eosinophil count Go to 21				
	☐ IgE level Go to 22				
1 In the 12 months immediately prior to commencing PBS-subsidised biological medicine treatment for severe asthma, the patient had:					
	a baseline blood eosinophil count ≥ 150 cells/microlitre while receiving treatment with OCS				
	Blood eosinophil count cells per microlitre				
	Date (DD MM YYYY)				
	or (not applicable to dupilumab 300 mg applications)				
	\Box a baseline blood eosinophil count \ge 300 cells/microlitre				
	Blood eosinophil count cells per microlitre				
	Date (DD MM YYYY)				
	Go to 24				
	In the 12 months immediately prior to commencing				
	 PBS-subsidised biological medicine treatment for severe asthma, the patient had: □ total serum human immunoglobulin E (IgE) ≥ 30 IU/mL with past or current evidence of atopy, documented by skin prick testing 				
	asthma, the patient had: ☐ total serum human immunoglobulin E (IgE) ≥ 30 IU/mL with past or current evidence of atopy, documented by skin prick				
3	 asthma, the patient had: total serum human immunoglobulin E (IgE) ≥ 30 IU/mL with past or current evidence of atopy, documented by skin prick testing or total serum human IgE ≥ 30 IU/mL with past or current evidence of atopy, documented by an in vitro measure of 				
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24	asthma, the patient had: total serum human immunoglobulin E (IgE) ≥ 30 IU/mL with past or current evidence of atopy, documented by skin prick testing or total serum human IgE ≥ 30 IU/mL with past or current evidence of atopy, documented by an in vitro measure of specific IgE Provide the patient's total serum human IgE (no older than 12 months immediately prior to commencing PBS-subsidised biological medicine treatment for severe asthma) IgE result IU/mL Date (DD MM YYYY) Comments in the relevant attachments need to be provided with this form.				

Personal information may be used by Services Australia, or 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).

More information about the way in which Services Australia manages personal information, including our privacy policy, can be found at **servicesaustralia.gov.au/privacypolicy**

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**

26 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)

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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
 or
- by post (signature required) to

Services Australia Complex Drugs Programs Reply Paid 9826 HOBART TAS 7001