

medicare



Ankylosing spondylitis – continuing authority application

When to use this form

Use this form to apply for **continuing** PBS-subsidised biological medicines (**originator** brands) for patients 18 years or over with ankylosing spondylitis.

Applications for **continuing** treatment with PBS-subsidised **biosimilar** brands of adalimumab, etanercept and infliximab are **Authority Required (STREAMLINED)** and do not require authority approval from Services Australia for the listed quantity and repeats.

Important information

Continuing authority applications for **originator** brands 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.

Under no circumstances will phone approvals be granted for ankylosing spondylitis **continuing** authority applications for **originator** brands.

Where the term 'biological medicine' appears, it refers to adalimumab, bimekizumab, certolizumab pegol, etanercept, golimumab, infliximab, ixekizumab, secukinumab, tofacitinib and upadacitinib.

A copy of the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) is provided for your convenience, but is not required to be submitted with this application.

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 **continuing** treatment with an **originator** brand.

The patient remains eligible to receive **continuing** treatment providing they continue to sustain a response to treatment.

Section 100 arrangements for infliximab i.v.

This item is 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.

This item is 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.

For more information

Go to servicesaustralia.gov.au/healthprofessionals

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Patient's details		Hospital details
1	Medicare card number Ref no. Or Department of Veterans' Affairs card number	8 Hospital name This hospital is a: public hospital
2	Dr	private hospital 9 Hospital provider number
	First given name	To qualify for PBS authority approval, the following conditions must be met.
3	Date of birth (DD MM YYYY) Patient's current weight kg	10 The patient is being treated by a: rheumatologist clinical immunologist with expertise in the management ankylosing spondylitis
Pr	escriber's details	11 This application is for: adalimumab
5	Prescriber number	bimekizumab certolizumab pegol etanercept golimumab
6	Family name First given name	infliximab i.v. infliximab s.c. ixekizumab secukinumab tofacitinib
7	Business phone number (including area code) Alternative phone number (including area code)	upadacitinib 12 Has the patient previously received this biological medicine (regardless of formulation) as their most recent course of PBS-subsidised treatment for this condition? No Yes Dates of the most recent treatment course From (DD MM YYYY) To (DD MM YYYY)

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The patient has demonstrated an adequate response to treatment evidenced by: a BASDAI score of Date of assessment (DD MM YYYY)	Prescriber's declaration You do not need to sign the declaration if you complete using Adobe Acrobat Reader and return this form through Professional Online Services (HPOS) at servicesaustralia.gov.au/hpos
and an erythrocyte sedimentation rate (ESR) level of mm/hr Date of test (DD MM YYYY) and/or	 I declare that: I am aware that this patient must meet the cri the current Schedule of Pharmaceutical Benef eligible for this medicine. I have informed the patient that their personal (including health information) will be disclosed Australia for the purposes of assessing and prauthority application. I have provided details of the proposed prescri
a C-reactive protein (CRP) level of mg/L Date of test (DD MM YYYY) Where only one marker (ESR or CRP) has been provided at baseline, the same marker must be used for assessment for all continuing applications. All measurements must be no more than 4 weeks old at the time of application. Checklist	the relevant attachments as specified in the Pharmaceutical Benefits Scheme restriction. the information I have provided in this form is a correct. I understand that: giving false or misleading information is a serion 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
The relevant attachments need to be provided with this form. Details of the proposed prescription(s).	Potential this form
Privacy notice	Returning this form Return this form, details of the proposed prescription(s) relevant attachments:
Personal information is protected by law (including the Privacy Act 1988) and is collected by Services Australia for the purposes of assessing and processing this authority application. Personal information may be used by Services Australia, or	online (no signature required), upload through HP(servicesaustralia.gov.au/hpos or by post (signature required) to

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

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Date (DD MM YYYY) (you must date this declaration)		
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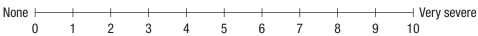
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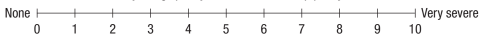
Bath Ankylosing Spondylitis Disease Activity Index (BASDAI)

Place a mark on each line below to indicate your answer to each question as it relates to your **past week**.





2 How would you describe the overall level of Ankylosing spondylitis neck, back or hip pain you have had?



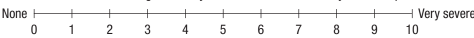
3 How would you describe the overall level of pain/swelling in joints other than your neck, back or hips that you have had?



4 How would you describe the overall level of discomfort you have had from any areas tender to touch or pressure?



5 How would you describe the overall level of morning stiffness you have had from the time you wake up?



6 How long does your morning stiffness last from the time you wake up?



Scoring the BASDAI

Measure each question from 'None' to the patient's mark in centimetres.

Add Q5 and Q6 and divide by 2 = A

Add Q1, Q2, Q3 and Q4 = B

Add A and B and divide by 5 = BASDAI score

BASDAI prepared by the Pharmaceutical Benefits Branch, Australian Government Department of Health and Aged Care, 15 July 2004. Reproduced and extracted from: Garrett, Sarah et al. (1994) A New Approach to Defining Disease Status in Ankylosing Spondylitis: The Bath Ankylosing Spondylitis Activity Index. Journal of Rheumatology, 21 (12), 2286–2291, with the permission of the copyright holder.