

# Non-radiographic axial spondyloarthritis – bimekizumab – 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 bimekizumab for patients with non-radiographic axial spondyloarthritis who have received non-PBS-subsidised treatment with bimekizumab for the same condition prior to **1 October 2024**.

# **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 non-radiographic axial spondyloarthritis **initial grandfather** authority applications.

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 **initial grandfather** treatment.

A patient may only qualify for PBS-subsidised treatment under this restriction once in a lifetime.

For **continuing** PBS-subsidised treatment, a grandfathered patient must qualify under the **continuing** treatment criteria.

After an authority application for **initial grandfather** treatment has been approved, applications for **continuing** treatment with bimekizumab 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.

# **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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Conditions and criteria



# **Online PBS Authorities** 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) Prescriber's details Prescriber number Family name First given name Business phone number (including area code)

Alternative phone number (including area code)

|    | qualify for PBS authority approval, the following conditions ust be met.   |
|----|--|
| 7  | The patient is being treated by a:  rheumatologist  clinical immunologist with expertise in the management of  |
|    | non-radiographic axial spondyloarthritis   |
| 8  | Has the patient previously received non-PBS-subsidised treatment with this biological medicine for this condition prior to <b>1 October 2024</b> ?  Yes \( \subseteq \text{No } align* |
| 9  | Has the patient responded inadequately to biological medicine on 4 occasions within the current treatment cycle?  Yes  No  |
| 10 | Prior to commencing non-PBS-subsidised treatment with this drug, was the condition diagnosed as non-radiographic axial spondyloarthritis, as defined by Assessment of Spondyloarthritis International Society (ASAS) criteria?  Yes No                                     |
| 11 | Prior to commencing non-PBS-subsidised treatment with this drug, was the condition radiographically evidenced on plain x-ray of Grade II bilateral sacroiliitis or Grade III or IV unilateral sacroiliitis?  Yes  No   |
| 12 | Prior to commencing non-PBS-subsidised treatment with this drug, the condition was:  sacroillitis with active inflammation on non-contrast Magnetic Resonance Imaging (MRI)  |
|    | and/or   |
|    | sacroiliitis with oedema on non-contrast MRI.  |
| 13 | Prior to commencing non-PBS-subsidised treatment with this drug, did the condition have presence of Bone Marrow Oedema (BMO) depicted as a hyperintense signal on a Short Tau Inversion Recovery (STIR) image (or equivalent)?  Yes  |
|    | No 🗔   |



|    | Prior to commencing non-PBS-subsidised treatment with this drug, did the condition have BMO depicted as a hypointense signal on a T1 weighted image (without gadolinium)?  Yes  No  Prior to commencing non-PBS-subsidised treatment with this drug for this condition, the patient had the following conditions: | Provide details of contraindications or intolerances to NSAID prior therapy, including the degree of toxicity.  For details of the toxicity criteria, go to servicesaustralia.gov.au/healthprofessionals  Intolerances must be of a severity to necessitate permanent treatment withdrawal. |
|----|---|---|
|    | Tick all that apply   |   |
|    | enthesitis (heel)   |   |
|    | uveitis   | 20 The patient's failure to achieve an adequate response to NSAID treatment and concomitant exercise program was demonstrated   |
|    | dactylitis  | by:   |
|    | psoriasis   | a BASDAI score of at least 4 on a 0–10 scale  |
|    | inflammatory bowel disease  | Baseline BASDAI score   |
|    | positive for Human Leukocyte Antigen B27 (HLA-B27).   |   |
| 16 | Prior to commencing non-PBS-subsidised treatment with this drug for this condition, did the patient have chronic lower back pain and stiffness for 3 or more months that was relieved by exercise but not rest?  Yes  | Date of assessment (DD MM YYYY)  and  an elevated C-reactive protein (CRP) > 10 mg/L  |
|    | No  | Baseline CRP level  |
| 17 | Prior to commencing non-PBS-subsidised treatment with   | Date of test (DD MM VVVV)   |
|    | this drug for this condition, did the patient fail to achieve an adequate response following treatment with at least 2 non-   | Date of test (DD MM YYYY)   |
|    | steroidal anti-inflammatory drugs (NSAIDs), whilst completing   | the requirement to demonstrate an elevated CRP could not  |
|    | an appropriate exercise program, for a total of 3 months?  Yes <b>Go to 18</b>  | be met due to   |
|    | No Go to 19   | treatment with prednisolone dosed at 7.5mg or   |
| 12 | Provide details of NSAID treatment  | higher daily (or equivalent)  |
|    | a) NSAID  | treatment with a parenteral steroid within the past month (intramuscular or intravenous   |
|    | ma  | methylprednisolone or equivalent) <b>or</b>   |
|    | Dose  | provide an acceptable reason the patient could not  |
|    | From (DD MM YYYY)   | demonstrate an elevated CRP level.  |
|    | To (DD MM YYYY)   |   |
|    | b) NSAID  | The baseline BASDAI score and CRP level must have been  |
|    | Dose mg   | determined at the completion of the 3-month NSAID and exercise trial, but prior to ceasing NSAID treatment.   |
|    | From (DD MM YYYY)   | The BASDAI and CRP results must have been <b>no more than</b> 4 weeks old at the time of initiating non-PBS-subsidised  |
|    | To (DD MM YYYY)   | treatment with this drug for this condition.  |
|    | If the NSAID dose was less than the maximum recommended dose in the relevant Therapeutic Goods Administration (TGA)   | These <b>baseline</b> results will need to be provided for all continuing applications to demonstrate the patient's   |
|    | approved Product Information, state the reason why.   | response.   |
|    |   | 21 Has the patient demonstrated an adequate response following at least 12 weeks of non-PBS-subsidised treatment with this drug for this condition?   |
|    |   | No U  |
|    |   | Not applicable as the patient has not had  12 weeks of treatment to   |
|    |   | demonstrate a response Go to 23   |

| 22        | The | patient's adequate response is evidenced by:                 |  |  |
|-----------|-----|--|--|--|
|           |     | a BASDAI score reduced by at least 2 from baseline           |  |  |
|           |     | Current BASDAI score   |  |  |
|           |     |  |  |  |
|           |     | Date of assessment (no more than 4 weeks old) (DD MM YYYY)   |  |  |
|           |     |  |  |  |
|           | and |  |  |  |
|           |     | a CRP level ≤ 10 mg/L  |  |  |
|           |     | Current CRP level  |  |  |
|           |     | mg/L   |  |  |
|           |     | Date of test (no more than 4 weeks old) (DD MM YYYY)         |  |  |
|           |     |  |  |  |
|           | or  |  |  |  |
|           |     | a CRP level reduced by at least 20% from baseline            |  |  |
|           |     | Current CRP level  |  |  |
|           |     | mg/L   |  |  |
|           |     | Date of test (no more than 4 weeks old) (DD MM YYYY)         |  |  |
|           |     |  |  |  |
|           | or  |  |  |  |
|           |     | CRP exempt at baseline                                       |  |  |
| Checklist |     |  |  |  |
| 23        | G   | The relevant attachments need to be provided with this form. |  |  |
|           | Ë   |  |  |  |
|           |     | Details of the proposed prescription(s).                     |  |  |
|           |     |  |  |  |

# **Privacy notice**

**24** 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 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

### 25 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.

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|--|--|--|--|--|
| • giving false or misleading information is a serious offence. |  |  |  |  |
| I have read, understood and agree to the above.                |  |  |  |  |
| Date (DD MM YYYY) (you <b>must</b> 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

٥r

by post (signature required) to

Services Australia Complex Drugs Programs Reply Paid 9826 HOBART TAS 7001



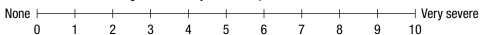
# medicare



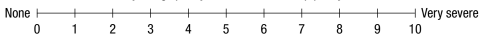
# Bath Ankylosing Spondylitis Disease Activity Index (BASDAI)

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

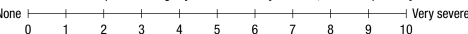




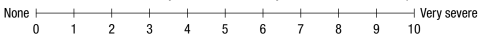
2 How would you describe the overall level of Ankylosing spondylitis neck, back or hip pain you have had?



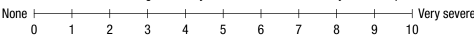
3 How would you describe the overall level of pain/swelling in joints other than your neck, back or hips that you have had?



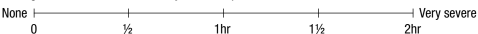
How would you describe the overall level of discomfort you have had from any areas tender to touch or pressure?



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 = Score

BASDAI prepared by the Pharmaceutical Benefits Branch, Australian Government Department of Health, 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.