

medicare



Psoriatic arthritis – initial authority application

When to use this form	Use this form to apply for initial PBS-subsidised biological medicines for patients 18 years or over with severe psoriatic arthritis.		
Important information	Initial applications to start PBS-subsidised treatment 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 psoriatic arthritis initial authority applications.		
	Where the term 'biological medicine' appears, it refers to adalimumab, bimekizumab, certolizumab pegol, etanercept, golimumab, guselkumab, infliximab, ixekizumab, risankizumab, secukinumab, tofacitinib, upadacitinib and ustekinumab. Patients are eligible for PBS-subsidised treatment with only one biological medicine at any time.		
	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 treatment.		
	After a written authority application for the first continuing treatment has been approved, subsequent continuing treatments with PBS-subsidised biosimilar brands of biological medicines are Authority Required (STREAMLINED) and do not require authority approval from Services Australia for the listed quantity and repeats.		
Section 100 arrangements	This item is available to a patient who is attending:		
for infliximab i.v.	• 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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PBS

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Patient's details			Hospital details		
1	Medicare card number	8	Hospital name		
	or		This hospital is a:		
	Department of Veterans' Affairs card number		public hospital		
			private hospital		
2	Dr 🗌 Mr 🗌 Mrs 🗌 Miss 🗌 Ms 🗌 Other 🦳	9	Hospital provider number		
	First given name	Co	nditions and criteria		
			o qualify for PBS authority approval, the following conditions ust be met.		
3	Date of birth (DD MM YYYY)	10	The patient, 18 years or over, is being treated by a:		
			rheumatologist		
4	Patient's weight kg		clinical immunologist with expertise in the management of psoriatic arthritis		
		11	This application is for:		
Pr	escriber's details		adalimumab		
_			bimekizumab		
5	Prescriber number		certolizumab pegol		
			etanercept		
6	Dr 🗌 Mr 🗌 Mrs 🗌 Miss 🗌 Ms 🗌 Other		golimumab		
U	Family name		guselkumab		
			infliximab i.v.		
	First given name		infliximab s.c. with i.v. loading (and an authority prescription for at least 2 i.v. doses at weeks 0 and 2 is attached)		
7	Dusiness phone number (including erec code)		ixekizumab		
1	Business phone number (including area code)		risankizumab		
			secukinumab		
	Alternative phone number (including area code)		tofacitinib		
			upadacitinib		
			ustekinumab		



The patient:	14 The patient has failed to achieve an adequate response to prior
has not received PBS-subsidised treatment with a	DMARD treatment demonstrated by:
biological medicine for this condition	an elevated erythrocyte sedimentation rate (ESR) > 25 mm/h
	Baseline ESR level mm/hr
has failed to achieve an adequate response following a minimum of 3 months treatment with:	Date of test (DD MM YYYY)
methotrexate, at a dose of at least 20 mg/week	
	and/or
From (DD MM YYYY)	an elevated C-reactive protein (CRP) > 15 mg/L
	Baseline CRP level mg/L
and	Date of test (DD MM YYYY)
sulfasalazine, at a dose of at least 2 g/day	
From (DD MM YYYY)	or
	the requirement to demonstrate an elevated ESR or CRP
To (DD MM YYYY)	could not be met due to
	treatment with prednisolone dosed at 7.5mg or higher
or	daily (or equivalent)
leflunomide, at a dose up to 20 mg/day	or
From (DD MM YYYY)	treatment with a parenteral steroid within the past
	month (intramuscular or intravenous
To (DD MM YYYY)	methylprednisolone or equivalent)
	or
applicable, provide details of contraindications or intolerand	provide an acceptable reason the patient could not
prior disease-modifying anti-rheumatic drugs (DMARD)	demonstrate an elevated ESR or CRP level
tment, including the degree of toxicity.	
r details of the toxicity criteria, go to ervicesaustralia.gov.au/healthprofessionals	and
tolerance must be of a severity to necessitate permanent	an active joint count of at least 20 active (swollen and
reatment withdrawal.	tender) joints
rior therapy contraindication or toxicity and grade	Baseline total active joint count
Aethotrexate Grad	e Date of assessment (DD MM YYYY)
ulfasalazine Grad	e or
	at least 4 active major joints from elbow, wrist, knee,
eflunomide Grad	e ankle, shoulder and/or hip
	Baseline major joint count
	Date of assessment (DD MM YYYY)
	The baseline joint count and FCD and/or CDD lovel must be
	The baseline joint count and ESR and/or CRP level must be determined at the completion of the 3-month DMARD trial,
	but prior to ceasing DMARD therapy.
	All measures must be no more than 4 weeks old at the time
	of initial application.
	Where only one marker (ESR or CRP) has been provided at baseline, the same marker must be used for assessment for
	all continuing applications.

Where a patient has at least 4 active major joints and less than 20 total active joints at baseline, assessment of the major joints only will be used for all continuing applications.

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Checklist

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The relevant attachments need to be provided with this form.

Details of the proposed prescription(s).

Privacy notice

16 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**

17 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