

Kaiser Permanente Health Plan of Mid-Atlantic States, Inc.
Ilumya (tildrakizumab-asmn) Prior Authorization (PA)
Pharmacy Benefits Prior Authorization Help Desk
Length of Authorization: 12 months

Instructions:

This form is used by Kaiser Permanente and/or participating providers for coverage of **Ilumya (tildrakizumab-asmn)**. Please complete and fax this form back to Kaiser Permanente within 24 hours [fax: <u>1-866-331-2104</u>]. If you have any questions or concerns, please call <u>1-866-331-2103</u>. **Requests will not be considered unless this form is complete. The KP-MAS Formulary can be found at: http://www.providers.kaiserpermanente.org/mas/formulary.html**

	1 – Patient Information	
Patient Name:	Kaiser Medical ID#:	Date of Birth:
	2 – Provider Information	
Provider Name:	Specialty:	Provider NPI:
Provider Address:		
Provider Phone #:	Provider Fax #:	
Please check the boxes that apply: □ Initial Request □ Continuation of	Therapy Request	
	3 – Pharmacy Information	
Pharmacy Name:	Pharmacy NPI:	
Pharmacy Phone #	Pharmacy Fax #:	
	4 – Drug Therapy Requested	
Drug 1: Name/Strength/Formulation	n:	
Drug 2: Name/Strength/Formulation	n:	
5,8.		-

5- Diagnosis/Clinical Criteria

	□ Plaque Psoriasis (PsO) □ Psoriatic Arthritis (PsA)		
	□ Othe	er:	
2.	Was there therapeutic failure on oral methotrexate? AND□ No □ Yes		
3.	Was th □ No □	nere therapeutic failure to one of the preferred agents? Yes	(e.g. Enbrel, Humira) AND
4.	 4. If this is being used for <u>plaque psoriasis</u> (PSO): a. Does the patient have moderate-to-severe plaque psoriasis for at least 6 months? AND □ No □ Yes 		riasis for at least 6 months? AND
	b.	Is there involvement of at least 10% of body surface a $\hfill \square$ No $\hfill \square$ Yes	rea (BSA)? OR
	C.	Is the Psoriasis Area and Severity Index (PASI) score 10 \Box No \Box Yes	or greater? OR
	 d. Incapacitation due to plaque location (e.g., head and neck, palms, soles or genitalia)? AND □ No □ Yes 		
	e.	Has the patient not responded adequately (or is not a agents (e.g., anthralin, coal tar preparations, corticost retinoic acid derivatives, and/or Vitamin D analogues) □ No □ Yes	eroids, emollients, immunosuppressives, keratolytics,
	f.	Has the patient not responded adequately (or is not a systemic agent (e.g. Immunosuppressives, retinoic aci \Box No \Box Yes	
phot		as the patient not responded adequately (or is not a candidate) to a 3 month minimum trial of hototherapy (e.g. Psoralens with UVA light (PUVA) OR UVB with coal tar or dithranol)? No \Box Yes	
		6 – Provider Sign-	Off
ditio	onal Info	ormation – Please provide any additional information t	
cert	ify that t	he information provided is accurate. Supporting document	ation is available for State audits.
ovide	er Signatı	ure:	Date:

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