

Prior Authorization Request Form Fax Back To: (866) 940-7328 Phone: (800) 310-6826

Specialty Medication Prior Authorization Cover Sheet

(This cover sheet should be submitted along with a Pharmacy Prior Authorization Medication Fax Request Form. Please refer to www.uhcprovider.com for medication fax request forms.)

Patient Information			
Patient's Name:			
Insurance ID:	Date of Birth:	Height: Weight:	
Address:		Apartment #:	
City:	State:	Zip Code:	
Phone Number:	Alternate Phone:	Sex: ☐ Male ☐ Female	
Provider Information			
Provider's Name:	Provider ID Number:		
Address:	City:	State: Zip Code:	
Suite Number:	Building Number:		
Phone Number:	Fax number:		
Provider's Specialty:			
Medication Information			
Medication:	Quantity:	ICD10 Code:	
Directions:	Diagnosis:	Refills:	
Physician Signature**:	•	Initial here if DAW:	
		y pharmacy with a prescription	
Physician Signature**: Physician Signature**: By signing above, the partner that can be used to facilitate the dispensing and Medication Instructions	d/or coordination of delivery for t	y pharmacy with a prescription he requested medication.	
Physician Signature**: Physician Signature**: By signing above, the partial that can be used to facilitate the dispensing and Medication Instructions Has the patient been instructed on how to Selection	d/or coordination of delivery for t	y pharmacy with a prescription he requested medication.	
Physician Signature**: Physician Signature**: By signing above, the puthat can be used to facilitate the dispensing and Medication Instructions Has the patient been instructed on how to Sells this medication a New Start?	d/or coordination of delivery for t	y pharmacy with a prescription he requested medication. Yes No Yes No	_
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Physician Signature**: By signing above, the puthat can be used to facilitate the dispensing and Medication Instructions Has the patient been instructed on how to Sells this medication a New Start? If continuation please provide the following: Is there documentation of positive clinical references attach any pertinent clinical inform Additional clinical information may be need.	d/or coordination of delivery for to delivery for the delivery for to delivery for the	y pharmacy with a prescription he requested medication. Yes No Yes No Date of Last Dose: / / Yes No upport stated diagnosis.	
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Physician Signature**: By signing above, the puthat can be used to facilitate the dispensing and Medication Instructions Has the patient been instructed on how to Sells this medication a New Start? If continuation please provide the following: Is there documentation of positive clinical reference attach any pertinent clinical inform Additional clinical information may be need previously tried and failed. Delivery Instructions Note: Delivery coordination requires a "Physical Provider Information" and "Patient Information" and "P	If-Administer? Initiation Date: / / esponse to current therapy? mation that would pertain to suded depending on your patient ician Signature" above and control information a	y pharmacy with a prescription he requested medication. Yes No Yes No Date of Last Dose: / / Yes No Ipport stated diagnosis. Is plan, including medication(s)	





PRIOR AUTHORIZATION REQUEST FORM

Please complete this entire form and fax it to: 866-940-7328. If you have questions, please call 800-310-6826.

This form contains multiple pages. Please complete all pages to avoid a delay in our decision.

Allow at least 24 hours for review.

Section A – Member Inform	nation						
First Name:	Last Name:				Member ID:		
Address:							
City:		State:			ZIP (Code:	
Phone:		DOB:			Aller	gies:	
Primary Insurance Information:							
Is the requested medication	n 🗆 New or 🗆 C	Continuation	on of Therapy? If o	continuation, lis	st star	t date:	
Is this patient currently hos		Yes □ No	If recently discha	arged, list disch	narge	date:	
Section B - Provider Inform	nation						
First Name:			Last Name:				M.D./D.O.
Address:			City:		State) :	ZIP code:
Phone:	Fax:		NPI #:		Spec	cialty:	
Office Contact Name / Fax a	ttention to:						
Section C - Medical Inform	ation						
Medication:					S	trength:	
Directions for use:					Q	uantity:	
Diagnosis (Please be specif	fic & provide as	much infor	mation as possible	e):	IC	D-10 COD	DE:
Is this member pregnant?		If yes	, what is this mem	nber's due date	?		
Section D - Previous Med	lication Trials					Bassa	
.)	lication Trials	If yes	, what is this men	Dates of The			on for failure /
Section D - Previous Med	lication Trials						
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	Community Plan	PR	OR AUTHORIZATION REQUEST FOR			
Member Fire	st name:	Member Last name:	Member DOB:			
	Clir	nical and Drug Specific I	nformation			
		ALL REQUESTS				
	Does the patient have one ☐ Active ankylosing spond ☐ Active psoriatic arthritis	e of the following diagnoses? (If ye	es, check which applies)			
	□ Moderate to severe chro	nic plaque psoriasis adenitis suppurativa (i.e., Hurley Sta	ge II or III)			
□ Yes □ No	□ No □ Moderately to severely active Crohn's disease □ Moderately to severely active polyarticular juvenile idiopathic arthritis					
	□ Moderately to severely a	ctive rheumatoid arthritis (RA)				
	☐ Moderately to severely a☐ Non-infectious uveitis (U					
□ Yes □ No		last 6 months, unless contraindic	of methotrexate at up to the maximally ated or clinically significant adverse effects			
□ Yes □ No	□ Biologic DMARD [e.g., E□ Janus kinase inhibitor [e.	nbrel (etanercept), Cimzia (certolizu	, ,			
		ANKYLOSING SPONDYLITIS	<i>/-</i>			
□ Yes □ No	ibuprofen, naproxen] at u	istory of failure to two NSAIDs (no p to maximally indicated doses, e ndicated or clinically significant a	on-steroidal anti-inflammatory drugs) [e.g., ach used for at least 4 weeks within the last			
		CHROHN'S DISEASE				
			wing conventional therapies at up to			
		· · · · · · · · · · · · · · · · · · ·	contraindicated or clinically significant			
- Vaa - Na	□ 6-mercaptopurine (Purin	ienced? (If yes, check which applied	s and complete Section D above)			
□ Yes □ No	☐ Azathioprine (Imuran)	etrioi)				
	□ Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)					
	□ Methotrexate (Rheumatr	ex, Trexall)				
□ Yes □ No	Has the patient lost respo		e.g., Remicade, Inflectra, Renflexis)?			
		HIDRADENITIS SUPPURATIV	A			
□ Yes □ No	rifampin) at up to maxima		al antibiotic (e.g., doxycycline, clindamycin, 3 months, unless contraindicated or es, complete Section D above)			
		PLAQUE PSORIASIS				
□ Yes □ No	Does the patient have gre or genital involvement, or		face area involvement, palmoplantar, facial,			
	months, unless contraind	istory of failure to any of the follo icated or clinically significant advantage and complete Section D above)	wing topical therapies within the last 3 rerse effects are experienced?			
□ Yes □ No	□ Coal tar	g., tacrolimus, pimecrolimus)				
	□ Corticosteroids (e.g., bet□ Tazarotene□ Vitamin D analogs (e.g.,	calcitriol calcipotriene)				
	- vitariiii b arialogo (c.g.,	oaroninoi, oaronpouriorioj				







Member First name:	Member Last name:	Member DOB:			
RHEUMATOID ARTHRITIS					
Does the patient have a history of failure to a 3 month trial of any non-biologic disease modifying anti- rheumatic drug (DMARD) [e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at up to maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced? (If yes, complete Section D above)					
	ULCERATIVE COLIT	IS .			
Does the patient have a history of failure to any of the following conventional therapies at up to maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced? (If yes, check which applies and complete Section D above) 6-mercaptopurine (Purinethol) Aminosalicylates (e.g., mesalamine, sulfasalazine) Azathioprine (Imuran) Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)					
	UVEITIS				
□ Yes □ No Is the patient's Intermediate Panuveitis Posterior	Uveitis classified as any of the following	ig? (If yes, check which applies)			
Does the patient have a history of failure to at least one corticosteroid (e.g., prednisolone, prednisone) at up to a maximally indicated dose within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced? (If yes, complete Section D above)					
☐ Yes ☐ No (e.g., methotre: within the last:	Does the patient have a history of failure to at least one systemic non-biologic immunosuppressant				
CONTINUATION OF THERAPY					
□ Yes □ No Is there documentation of positive clinical response to Humira therapy?					
		_			

Physician Signature: ______ Date: _____

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