## HUMIRA (ADALIMUMAB) [PREFERRED] PRIOR AUTHORIZATION FORM





(form effective 1/9/2023)

Fax to PerformRx<sup>SM</sup> at **1-888-981-5202**, or to speak to a representative call **1-866-610-2774**.

PRIOR AUTHORIZATION REQUEST INFO	PMATION							
Contact's phone number:		LTC facilit	y contact/phon					
PATIENT INFORMATION								
Patient name:	Patient ID #:					DOB:		
Street address:			Apt. #: City/state/zip:					
		7.94		only/ oracle	,			
PRESCRIBER INFORMATION			Chaolalhu					
scriber name:			Specialty:  MA Provider ID #					
State license #:	NPI:							
			e #: City/state/zip:					
Phone:			Fax:					
CLINICAL INFORMATION								
Product requested: ☐ Humira ☐ Humira CF ☐ other	:							
Strength and formulation/packaging (i.e., syringe, pen, sta	rter pack, etc.):							
Directions:				Qty:	Re	efills:	Patient's weight:	lbs/kg
Diagnosis (submit documentation):							Dx code ( <u>required</u> ):	
PHARMACY INFORMATION (Prescriber t	o identify the ph	armacy	that is to c	lisnense i	the med	dication).		
,	Patient's Preferred Ph			лэрспэс	tire me	alcation).		
Pharmacy Phone #: Pharmacy Fax #:								
$\hfill \square$ I acknowledge that the patient agrees with the pharmac	y chosen for delivery o	f this medic	cation.					
1. All diagnoses: Check all that apply to the patient and su screened for hepatitis B (anti-HBs, HBsAg, and anti-H 2. All diagnoses: Is Humira being prescribed by or in consumer Yes — List specialty:  3. Ankylosing spondylitis: Does the patient have a history Yes — List medications tried:  No — Provide explanation:  4. Crohn's disease: Does at least one of the following app moderate to severe Crohn's disease and one of the fill failed to achieve remission with or has a contraining has one or more high-risk or poor prognostic feature has achieved remission with the requested medications.  5. Ulcerative colitis: Check all that apply to the patient.  mild UC that is associated with multiple poor prognosim moderate-to-severe UC  failed to achieve remission with or has a contraindical failed to achieve remission with or has a contraindical failed to achieve remission with or has a contraindical failed to achieve remission with or has a contraindical failed to achieve remission with or has a contraindical failed to achieve remission or the acceptable to the second of the failed to achieve remission with or has a contraindical failed to achieve remission or the acceptable to the second of the failed to achieve remission with or has a contraindical failed to achieve remission or the acceptable to the failed to achieve remission or the acceptable to the failed to achieve remission or the acceptable to the failed to achieve remission or	IBc)  screened for station with an appropriate of trial and failure of a by to the patient? Sollowing: dication or intolerance to company the stic factors	tuberculosis riate special two-week to an induction e requested an induction	trial of continue	corticosteroid lators maintenanc	ds			
☐ failed to maintain remission or has a contraindicatior ☐ has achieved remission with the requested medication	on and will be using the	e requested	medication as	maintenanc				
6. Rheumatoid arthritis: Does the patient have a history of tri ☐ Yes — List medications tried or explain contraindication	on:	cation, or in	tolerance of at I				otrexate or another non-bi	iologic DMARD? No
7. Juvenile idiopathic arthritis (JIA): Check all that apply to the therapeutic failure of a three-month trial of a conven contraindication or intolerance to non-biologic DMAR systemic JIA with active systemic features one or more risk factors for disease severity involvement of high-risk joints (e.g., cervical spine, high disease activity is at high risk of disabling joint damage active sacrolitis and/or enthesitis and has tried and the sacrolitis and/or enthesitis and the sacrolitis and th	tional non-biologic DM/ Ds; provide explanation ip, wrist)	n:						

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INITIAL REQUESTS (Complete questions applicable to patient's diagnosis):						
8. Psoriatic arthritis: Does at least one of the following apply to the patient?						
a axial disease, dactylitis, and/or enthesitis						
has tried and failed methotrexate or other DMARD for at least 8 weeks; list medications tried or explain contraindication:						
□ severe disease						
concomitant moderate-to-severe nail disease						
□ concomitant active inflammatory bowel disease						
9. <u>Chronic psoriasis:</u> Check all that apply to the patient.						
☐ at least 3% of body surface area (BSA) is affected						
☐ critical areas of the body are involved (such as face, palms, soles, and/or genitals)						
☐ significant disability or impairment of physical, mental, or psychosocial functioning						
☐ moderate to severe nail disease						
☐ history of therapeutic failure, contraindication or intolerance to (check all that apply):						
🗆 4-week trial of topical steroids or 8-week trial of other topical therapy; list medications tried or explain contraindication:						
□ 3-month trial of conventional systemic therapy; list medications tried or explain contraindication:						
□ phototherapy						
10. <u>Uveitis:</u> Check all of the following that apply to the patient and submit documentation for each.						
☐ has a diagnosis of uveitis associated with juvenile idiopathic arthritis or Behçet's disease						
☐ has steroid-dependent uveitis (i.e., requires ≥ prednisone 7.5 mg daily [or equivalent]) with plan to taper or discontinue systemic steroids						
☐ has a documented history of trial and failure, contraindication, or intolerance of systemic immunosuppressives or corticosteroids (systemi	c, topical, intraocular, or periocular)					
List medications tried:						
11. <u>Hidradenitis suppurativa (HS):</u> Check all that apply to the patient.						
☐ Hurley stage II disease						
☐ Hurley stage III disease						
☐ history of therapeutic failure, contraindication, or intolerance to:						
☐ 3-month trial of topical clindamycin						
☐ adequate trial of a systemic antibiotic						
☐ Is a candidate for or has history of surgical intervention for HS						
12. All other diagnoses: Submit documentation supporting the use of Humira for the patient's diagnosis and other treatments tried.						
RENEWAL REQUESTS						
1. Since starting Humira, has the patient experienced improvement in disease activity and/or level of functioning?  ☐ Yes ☐ No						
2. Is Humira being prescribed by or in consultation with an appropriate specialist?						
Yes - list specialty:						
PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION						
Prescriber signature:	Date:					

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