CYTOKINE AND CAM ANTAGONISTS PRIOR AUTHORIZATION FORM





(form effective 1/8/2024)

Fax to PerformRx[™] at **1-855-851-4058**, or to speak to a representative call **1-866-907-7088**.

PRIOR AUTHORIZ	ATION REQU	EST INFORMATION					
□ New request □ Renewal request Total # of pages:							
Name of office contact: Contact's			Contact's p	phone number: LTC fa		LTC fac	cility contact/phone:
PATIENT INFORM	ATION						
Patient name:				Patient ID #:			DOB:
Street address:			· ·				
Apt #:	City/state/zip:				Phone:		
PRESCRIBER INFO	ORMATION						
Prescriber name:							
Specialty:				NPI:			State license #:
Street address:							
Suite #:	City/state/zip:						
Phone:	Phone:			Fax:			
CLINICAL INFORM	1ATION						
Medication requested: Preferred Medications: Actemra (tocilizuma Actemra (tocilizuma Adalimumab-fkjp(C) Adalimumab-fkjp(C) Syringe Avsola (infliximab-a Enbrel (etanercept) Enbrel (etanercept) Enbrel (etanercept) Hadlima (adalimum 50 mg/ml Pushtouc Hadlima(CF) (adalin 100 mg/ml Syringe Humira (adalimumal)	b) Vial F) 50 mg/ml Pen F) 50 mg/ml Mini Cartridge Sureclick Pen Syringe Vial ab-bwwd) ch ab-bwwd) umab-bwwd) umab-bwwd) b) 50 mg/ml Pen	 Humira (adalimumab) 50 mg/ml Syringe Humira(CF) (adalimumab) F Humira(CF) (adalimumab) S Infliximab Vial (Janssen's unbranded infliximab) Kineret (anakinra) Syringe Orencia (abatacept) Clickje Orencia (abatacept) Vial Otezla (apremilast) Tablet Simponi (golimumab) Pen Simponi (golimumab) Syringe Xeljanz (tofacitinib) Tablet Xeljanz XR (tofacitinib) Tablet Yusimry(CF) (adalimumab-a 50 mg/ml Pen 	Syringe It Ige ctor let aqvh)	 Actemr Adalim Adalim Syring Amjevii 50 mg Arcalys Cimzia Cosent Cyltezc 50 mg Cyltezc 50 mg Cyltezc 50 mg Cyltezc 50 mg Hulio(C 50 mg Hulio(C 50 mg Hulio(C 50 mg Hulio(C 50 mg Clintai Hulio(C 50 mg Clintai Hulio(C 50 mg Hulio(C 50 mg Hulio(C 50 mg Hulio(C 50 mg Idacio(50 mg Ilaris (c 	d Medications: a (tocilizumab) Actpen umab-adaz(CF) 100 mg/ml umab-adaz(CF) 100 mg/ml umab-adaz(CF) 100 mg/ml e ta(CF) (adalimumab-atto) /ml Autoinjector ta(CF) (adalimumab-atto) /ml Syringe t (rilonacept) Vial (certolizumab pegol) Syring yx (secukinumab) Pen yx (secukinumab) Pen yx (secukinumab) Pen yx (secukinumab) Syringe (CF) (adalimumab-adbm) /ml Pen (CF) (adalimumab-adbm) /ml Syringe z(CF) (adalimumab-fkjp) /ml Syringe z(CF) (adalimumab-adaz) g/ml Syringe anakinumab) Vial	ge	 Ilumya (tildrakizumab) Syringe Inflectra (infliximab-dyyb) Vial Kevzara (sarilumab) Pen Kevzara (sarilumab) Syringe Litfulo (ritlecitinib) Capsule Olumiant (baricitinib) Tablet Orencia (abatacept) Syringe Remicade (infliximab) Vial Renflexis (infliximab-abda) Vial Rinvoq ER (upadacitinib) Tablet Siliq (brodalumab) Syringe Simponi Aria (golimumab) Vial Skyrizi (risankizumab) On-Body Injector Skyrizi (risankizumab) Pen Skyrizi (risankizumab) Yial Sotyktu (deucravacitinib) Tablet Stelara (ustekinumab) Vial Stelara (ustekinumab) Vial Stelara (ustekinumab) Vial Stelara (ustekinumab) Syringe Xeljanz (tofacitinib) Solution Yuflyma(CF) (adalimumab-aaty) 100 mg/ml Autoinjector Yuflyma(CF) (adalimumab-aaty) 100 mg/ml Syringe
Quantity per fill: Refills:				Quantity per fill: Refills:			
Directions:				Directions:			
Diagnosis (submit documentation):							
Is the beneficiary currently being treated with the requested medication?				Dx code (required): Beneficiary weight: Press – date of last dose: Submit documentation.			
				No			
Is the requested medication prescribed by or in consultation with a specialist (e.g., rheumatologist, dermatologist, gastroenterologist, etc)?			g.,	□ Yes □ No If prescriber is not a specialist, submit documentation of consultation.			

	ARMACY INFORMATION (Prescriber to identify the pharmacy that is to dispense the medication): er to: Patient's Home Physician's Office Patient's Preferred Pharmacy Name:						
VPI#:							
harn	nacy Phone #: Pharmacy Fax #:						
∃la	cknowledge that the patient agrees with the pharmacy chosen for delivery of this medication.						
	Complete all sections that apply to the beneficiary and this request. Check all that apply and <u>submit documentation</u> for each item.						
	TIAL REQUESTS						
Drug 1.	Requested drug is NON-PREFERRED: Tried and failed or has a contraindication or intolerance to the preferred drugs in this class approved or medically accepted for the beneficiary's condition. List preferred medications tried:						
2.	Requested drug is OTEZLA (apremilast) or SILIQ (brodalumab): Was evaluated for history of prior suicide attempt, bipolar disorder, or major depressive disorder						
3.	 Requested drug is an oral JAK inhibitor (eg, Olumiant [baricitinib], Rinvoq [upadacitinib], Xeljanz [tofacitinib]): Tried and failed at least one TNF blocker or another biologic as recommended in the JAK inhibitor's package labeling Has a contraindication or an intolerance to TNF blockers or other biologics as recommended in the JAK inhibitor's package labeling 						
-	nosis <u>ALL diagnoses:</u> □ Screened for hepatitis B virus infection (surface antigen, surface antibody, and core antibody) □ Screened for tuberculosis						
2.	Adult-onset Still's disease: Has predominantly systemic disease: Has steroid-dependent disease with intent to decrease or discontinue dose of steroid with use of Cytokine and CAM Antagonist Tried and failed or has a contraindication or an intolerance to systemic glucocorticoids Has predominantly joint disease: Tried and failed or has a contraindication or an intolerance to conventional DMARDs (eg, MTX)						
3.	Alopecia areata: Has alopecia universalis Has >50% scalp involvement or alopecia totalis Has alopecia that causes significant disability or impaired physical, mental, or psychosocial functioning Has a current episode of alopecia areata that has lasted at least 6 months						
4.	Ankylosing spondylitis & non-radiographic axial spondyloarthritis: Tried and failed a 2-week trial of or has a contraindication or an intolerance to 2 different oral NSAIDs						
5.	Behçet's syndrome: □ Has a diagnosis of Behçet's syndrome according to current consensus guidelines □ Has recurrent oral ulcers associated with Behçet's syndrome □ Tried and failed or has a contraindication or an intolerance to a topical corticosteroid (eg, triamcinolone dental paste) □ Tried and failed a 3-month trial of or has a contraindication or an intolerance to colchicine at maximally tolerated doses						
6.	Crohn's disease: Has moderate-to-severe disease Has disease that is associated with high-risk or poor prognostic features Tried and failed to <u>achieve remission</u> with or has a contraindication or an intolerance to an induction course of corticosteroids Tried and failed to <u>maintain remission</u> with or has a contraindication or an intolerance to conventional immunomodulators (e.g., AZA, 6-MP, MTX)						
7.	Familial Mediterranean fever:						
8.	Gout flare: Tried and failed or has a contraindication or an intolerance to NSAIDs at maximally tolerated doses Tried and failed or has a contraindication or an intolerance to colchicine at maximally tolerated doses Tried and failed or has a contraindication or an intolerance to colchicine at maximally tolerated doses Tried and failed or has a contraindication or an intolerance to corticosteroids Has a medical reason why repeated courses of corticosteroids are not appropriate Output Description: Description:						
9.	Giant cell arteritis: Tried and failed or has a contraindication or an intolerance to systemic glucocorticoids Is at high risk for glucocorticoid-related complications Has steroid-dependent disease with intent to decrease or discontinue dose of steroid with use of Cytokine and CAM Antagonist						
10.	Hidradenitis suppurativa (HS): Has Hurley stage II or stage III disease Is a candidate for or has a history of surgical intervention for HS Tried and failed a 3-month trial of or has a contraindication or an intolerance to topical clindamycin Tried and failed or has a contraindication or an intolerance to systemic antibiotics (e.g., doxycycline, minocycline, tetracycline, clindamycin)						

INITIAL REQUESTS (continued)

11. Juvenile idiopathic arthritis:

- □ Has systemic disease with active systemic features
- □ Has disease associated with any of the following:
 - Positive anti-CCP antibodies
 - Positive rheumatoid factor
 - \Box Presence of joint damage
 - □ At high risk of disabling joint damage
 - □ High disease activity
 - □ Involvement of high-risk joints (cervical spine, hip, wrist)
- □ Tried and failed a 3-month trial of or has a contraindication or an intolerance to conventional DMARDs (e.g., MTX)
- □ Has active sacroiliitis and/or enthesitis:
- □ Tried and failed a 2-week trial of or has a contraindication or an intolerance to oral NSAIDs

12. Plaque psoriasis:

- \Box Has a BSA of \geq 3% that is affected
- □ Has involvement of critical areas of the body (eg, skin folds, face, genitals)
- □ Has psoriasis that causes significant disability or impaired physical, mental, or psychosocial functioning
- □ Has moderate-to-severe nail disease
- □ Tried and failed a 4-week trial of or has a contraindication or an intolerance to topical corticosteroids
- Tried and failed an 8-week trial of or has a contraindication or an intolerance to non-steroid topical medications (e.g., anthralin, calcineurin inhibitor, tazarotene, etc)

13. Polymyalgia rheumatica:

- Tried and failed or has a contraindication or an intolerance to systemic glucocorticoids
- □ Has steroid-dependent disease with intent to decrease or discontinue dose of steroid with use of Cytokine and CAM Antagonist

14. Psoriatic arthritis:

Tried and failed an 8-week trial of or has a contraindication or an intolerance to conventional DMARDs (e.g., AZA, leflunomide, MTX, SSZ)

- □ Has predominantly axial disease, dactylitis, and/or enthesitis
- \Box Has severe disease
- $\hfill\square$ Has comorbid moderate-to-severe nail psoriasis
- □ Has comorbid active inflammatory bowel disease

15. Rheumatoid arthritis:

□ Tried and failed a 3-month trial of or has a contraindication or an intolerance to conventional DMARDs (e.g., AZA, leflunomide, MTX, etc.)

16. Sarcoidosis:

□ Tried and failed or has a contraindication or an intolerance to systemic glucocorticoids

- \Box Has steroid-dependent disease
- Tried and failed or has a contraindication or an intolerance to a conventional DMARD (e.g., AZA, leflunomide, MTX, mycophenolate)

17. Ulcerative colitis:

- Has moderate-to-severe disease
- □ Has disease associated with multiple poor prognostic factors
- □ Tried and failed to achieve remission with or has a contraindication or an intolerance to an induction course of corticosteroids
- Tried and failed to maintain remission with or has a contraindication or an intolerance to conventional immunomodulators (e.g., AZA, cyclosporine, 6-MP, MTX)

18. Uveitis (non-infectious):

- □ Has comorbid juvenile idiopathic arthritis
- □ Has comorbid Behçet's syndrome
- □ Has steroid-dependent disease with intent to decrease or discontinue dose of steroid with use of Cytokine and CAM Antagonist
- □ Tried and failed or has a contraindication or an intolerance to systemic, topical, intraocular, or periocular corticosteroids
- Tried and failed or has a contraindication or an intolerance to conventional systemic immunosuppressives (e.g., AZA, MTX, MMF, etc)

19. Spevigo (spesolimab) for treatment of generalized pustular psoriasis (GPP) flares:

□ Has received a single dose of Spevigo (spesolimab) for current GPP flare:

- □ Continues to experience moderate to severe GPP flare symptoms since the previous dose
- □ Has not received a dose of Spevigo (spesolimab) for current GPP flare:
- □ Is experiencing a moderate to severe GPP flare that warrants rapid stabilization or improvement

20. Other diagnosis:

□ List other treatments tried (including start/stop dates, dose, outcomes):

RENEWAL REQUESTS

- Experienced an improvement in disease severity or level of functioning since starting therapy with the requested medication
- □ Is prescribed an increased dose or more frequent administration of the requested medication that is supported by peer-reviewed medical literature or national treatment guidelines □ Requested drug is OTEZLA (apremilast) or SILIQ (brodalumab):
- □ Was recently reevaluated for behavioral and mood changes

PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION							
Prescriber signature:	Date:						

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