## CYTOKINE AND CAM ANTAGONISTS PRIOR AUTHORIZATION FORM





(form effective 1/6/2025)

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

PRIOR AUTHORIZATION REQUI	EST INFORMATION					
☐ New request ☐ Renewal request	Total # of pages:					
Name of office contact:		Contact's	Contact's phone number:		LTC facility contact/phone:	
PATIENT INFORMATION						
Patient name:			Patient ID #:			DOB:
Street address:						
Apt #: City/state/zip:				Phone:		
PRESCRIBER INFORMATION						
Prescriber name:						
Specialty:			NPI:			State license #:
Street address:						
Suite #: City/state/zip:						
Phone:			Fax:			
CLINICAL INFORMATION						
Medication requested:						
Preferred Medication:  Adalimumab-aacf 50 mg/ml Pen Adalimumab-aacf 50 mg/ml Syringe Adalimumab-adaz(CF) 100 mg/ml Pen Adalimumab-adaz(CF) 100 mg/ml Syringe Adalimumab-adbm(CF) 50 mg/ml Pen (Boehringer Ingelheim) Adalimumab-adbm(CF) 50 mg/ml Syringe (Boehringer Ingelheim) Adalimumab-adbm(CF) 100 mg/ml Pen (Boehringer Ingelheim) Adalimumab-adbm(CF) 100 mg/ml Syringe (Boehringer Ingelheim) Adalimumab-adbm(CF) 50 mg/ml Syringe Adalimumab-fkjp(CF) 50 mg/ml Syringe Adalimumab-fkjp(CF) 50 mg/ml Syringe Amjevita(CF) (adalimumab-atto) 100 mg/ml Autoinjector	□ Amjevita(CF) (adalimumab-atto 100 mg/ml Syringe     □ Avsola (infliximab-axxq) Vial     □ Enbrel (etanercept) Mini Cartric     □ Enbrel (etanercept) Syringe     □ Enbrel (etanercept) Syringe     □ Enbrel (etanercept) Vial     □ Hadlima (adalimumab-bwwd)     50 mg/ml Pushtouch     □ Hadlima (adalimumab-bwwd)     50 mg/ml Syringe     □ Hadlima(CF) (adalimumab-bww 100 mg/ml Pushtouch     □ Hadlima(CF) (adalimumab-bww 100 mg/ml Syringe     □ Humira (adalimumab) 50 mg/ml	dge den vvd)	☐ Humira(ĈF☐ Humira(ĈF☐ Syringe☐ Infliximab☐ infliximab☐ Kineret (ar☐ Orencia (ac☐ Orencia (ac☐ Otezla (ap☐ Simlandi(C☐ 100 mg/m☐ Simponi (g☐ Simponi	akinra) Syringe batacept) Clickjet batacept) Vial emilast) Tablet FF) (adalimumab-ryvk) I Autoinjector olimumab) Pen olimumab) Syringe ankizumab-rzaa)	l Pen I	□ Skyrizi (risankizumab-rzaa) Pen □ Skyrizi (risankizumab-rzaa) Syringe □ Skyrizi (risankizumab-rzaa) Vial □ Taltz (ixekizumab) Autoinjector □ Taltz (ixekizumab) Syringe □ Tyenne (tocilizumab-aazg) Autoinjector □ Tyenne (tocilizumab-aazg) Vial □ Tyenne (tocilizumab-aazg) Vial □ Xeljanz (tofacitinib) Solution □ Xeljanz (tofacitinib) Tablet □ Xeljanz XR (tofacitinib) Tablet □ Yusimry(CF) (adalimumab-aqvh) 50 mg/ml Pen
Medication requested:  Non-Preferred Medication:  Abrilada(CF) (adalimumab-afzb) 50 mg/ml Pen  Abrilada(CF) (adalimumab-afzb) 50 mg/ml Syringe  Actemra (tocilizumab) Actpen  Actemra (tocilizumab) Syringe  Actemra (tocilizumab) Yial  Adalimumab-aaty(CF) 100 mg/ml Syringe  Adalimumab-aaty(CF) 50 mg/ml Pen (all labelers except Boehringer Ingelheim)  Adalimumab-adbm(CF) 50 mg/ml Syringe (all labelers except Boehringer Ingelheim)  Adalimumab-adbm(CF) 100 mg/ml Pen (all labelers except Boehringer Ingelheim)  Adalimumab-adbm(CF) 100 mg/ml Pen (all labelers except Boehringer Ingelheim)  Adalimumab-radbm(CF) 100 mg/ml Pen (all labelers except Boehringer Ingelheim)  Adalimumab-ryvk(CF) 100 mg/ml Syringe Ingelheim)  Adalimumab-ryvk(CF) 100 mg/ml Syringe  Adalimumab-ryvk(CF) 100 mg/ml Syringe  Amjevita(CF) (adalimumab-atto) 50 mg/ml Autoinjector	Amjevita(CF) (adalimumab-atto 50 mg/ml Syringe Arcalyst (rilonacept) Vial Bimzelx (bimekizumab-bkzx) A Bimzelx (bimekizumab-bkzx) S Cimzia (certolizumab pegol) Sy Cosentyx (secukinumab) Sensc Cosentyx (secukinumab) Vial Cytlezo(CF) (adalimumab-adbn 50 mg/ml Syringe Cytlezo(CF) (adalimumab-adbn 100 mg/ml Pen Cytlezo(CF) (adalimumab-adbn 100 mg/ml Pen Cytlezo(CF) (adalimumab-adbn 100 mg/ml Syringe Entyvio (vedolizumab) Pen Entyvio (vedolizumab) Pen Entyvio (vedolizumab) Pen Hulio(CF) (adalimumab-fkjp) 50 mg/ml Pen	utoinjector tyringe rringe oready Pen ge uady Pen n)	100 mg/m     Hyrimoz(C     100 mg/m     Idacio(CF)     50 mg/mI     Idacio(CF)     50 mg/mI     Ilaris (cana     Ilumya (tila     Inflectra (ii     Kevzara (s     Itifulo (rith)     Omvoh (m     Omvoh (m     Omvoh (m     Orencia (a     Renflexis (a     Rinvoq ER	F) (adalimumab-adaz) I Syringe (adalimumab-aacf) Pen (adalimumab-aacf)		□ Simponi Aria (golimumab) Vial □ Sotyktu (deucravacitinib) Tablet □ Spevigo (spesolimab-sbzo) Syringe □ Spevigo (spesolimab-sbzo) Vial □ Stelara (ustekinumab) Syringe □ Stelara (ustekinumab) Vial □ Tofidence (tocilizumab-bavi) Vial □ Tremfya (guselkumab) Autoinjector □ Tremfya (guselkumab) Syringe □ Yuflyma(CF) (adalimumab-aaty) 100 mg/ml Autoinjector □ Yuflyma(CF) (adalimumab-aaty) 100 mg/ml Syringe □ Zymfentra (infliximab-dyyb) Pen □ Zymfentra (infliximab-dyyb) Syringe



CLI	CLINICAL INFORMATION						
STARTER PACK requested (strength/formulation):		MAINTENANCE product/packaging requested (strength/formulation):					
Quan	tity per fill:	Refills:	Quantity per fill:	Refills:			
Direc	tions:		Directions:				
Diagr	nosis (submit documentation):		Dx code (required):	Beneficiary weight:			
Is the beneficiary currently being treated with the requested medication?			☐ Yes – date of last dose: ☐ No	Submit documentation.			
Is the requested medication prescribed by or in consultation with a specialist (e.g., rheumatologist, dermatologist, gastroenterologist, etc)?			☐ Yes ☐ No If prescriber is not a specialist, submit documentation of consultation.				
PH/	ARMACY INFORMATION (Pr	escriber to identify the pharmacy	that is to dispense the medication	n):			
	er to:  Patient's Home Physician's	s Office    Patient's Preferred Pharmacy Nar	ne:				
NPI#:	nacy Phone #:		Pharmacy Fax #:				
		the pharmacy chosen for delivery of this medic	-				
		Complete all sections that apply to Check all that apply and <u>submi</u>	o the beneficiary and this request. It documentation for each item.				
INI	TIAL REQUESTS						
Drug							
1.	<ul> <li>1. Requested drug is NON-PREFERRED:         <ul> <li>□ 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.</li> <li>List preferred medications tried:</li> </ul> </li> </ul>						
2.	2. Requested drug is BIMZELX (bimekizumab), OTEZLA (apremilast), or SILIQ (brodalumab):  ☐ Was evaluated for history of prior suicide attempt, bipolar disorder, or major depressive disorder						
3.	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						
Diag	nosis						
1.	<ul> <li>ALL diagnoses:</li> <li>□ Screened for hepatitis B virus infection (surface antigen, surface antibody, and core antibody) (if recommended in the FDA-approved package labeling)</li> <li>□ Screened for tuberculosis (if recommended in the FDA-approved package labeling)</li> </ul>						
2.	2. Adult-onset Still's disease:  Has predominantly systemic 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 (eq. MTX)						
3.	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.							
5.	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.		nigh-risk or poor prognostic features with or has a contraindication or an intolerance n with or has a contraindication or an intolerance		A, 6-MP, MTX)			
7.	Familial Mediterranean fever:  ☐ Tried and failed a 3-month trial of or	has a contraindication or an intolerance to colc	hicine at maximally tolerated doses				



## INITIAL REQUESTS (continued)

8.	Generalized pustular psoriasis (GPP) flares:  Request is for Spevigo (spesolimab) intravenous:  Is being treated for a GPP flare  One of the following:  Beneficiary has received a single dose of spesolimab for the current GPP flare AND:  Continues to experience moderate to severe GPP flare symptoms since the previous dose  Beneficiary has not received a dose of spesolimab for the current GPP flare AND:  Sexperiencing a moderate to severe GPP flare that warrants rapid stabilization or improvement  Request is for Spevigo (spesolimab) subcutaneous:  Has a history of at least one GPP flare  Is using subcutaneous spesolimab for the prevention of GPP flares
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.	Gout flare:  ☐ Tried and failed maximally tolerated doses of or has a contraindication or an intolerance to NSAIDs ☐ Tried and failed maximally tolerated doses of or has a contraindication or an intolerance to colchicine ☐ Tried and failed maximally tolerated doses of or has a contraindication or an intolerance to corticosteroids ☐ Has a medical reason why repeated courses of corticosteroids are not appropriate
	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)
	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  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 sacroillitis and/or enthesitis:  Tried and failed a 2-week trial of or has a contraindication or an intolerance to oral NSAIDs
	Plaque psoriasis:  ☐ Has a BSA of ≥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)
14.	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
	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   Has severe disease   Has comorbid moderate-to-severe nail psoriasis   Has comorbid active inflammatory bowel disease
16.	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.)
	Sarcoidosis:  □ Tried and failed or has a contraindication or an intolerance to systemic glucocorticoids □ Has steroid-dependent disease □ Tried and failed or has a contraindication or an intolerance to a conventional DMARD (e.g., AZA, leflunomide, MTX, mycophenolate)



INITIAL REQUESTS (continued)
18. <u>Ulcerative colitis:</u> ☐ Has moderate-to-severe disease ☐ Has disease associated with multiple poor prognostic factors ☐ 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, cyclosporine, 6-MP, MTX)
<ul> <li>19. Uveitis (non-infectious):         <ul> <li>Has comorbid juvenile idiopathic arthritis</li> <li>Has comorbid Behçet's syndrome</li> <li>Has steroid-dependent disease with intent to decrease or discontinue dose of steroid with use of Cytokine and CAM Antagonist</li> <li>Tried and failed or has a contraindication or an intolerance to systemic, topical, intraocular, or periocular corticosteroids</li> <li>Tried and failed or has a contraindication or an intolerance to conventional systemic immunosuppressives (e.g., AZA, MTX, MMF, etc.)</li> </ul> </li> <li>20. Other diagnosis:         <ul> <li>List other treatments tried (including start/stop dates, dose, outcomes):</li> </ul> </li> </ul>
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 BIMZELX (bimekizumab), OTEZLA (apremilast), or SILIQ (brodalumab): □ Was recently reevaluated for behavioral and mood changes
PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION

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Date:

Prescriber signature:

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