CYTOKINE AND CAM ANTAGONISTS PRIOR AUTHORIZATION FORM





(form effective 1/8/2024)

Fax to PerformRxSM at **1-888-981-5202**, or to speak to a representative call **1-866-610-2774**.

| PRIOR AUTHORIZ | ATION REQUI | EST INFORMATION | | | | | |
|--|----------------------|--|--|---|-----------------------|--|----------------------|
| ☐ New request ☐ Rei | newal request | Total # of pages: | | | | | |
| Name of office contact: Contact's | | | Contact's phor | phone number: LTC fa | | LTC fac | ility contact/phone: |
| PATIENT INFORMA | ATION | | | | | | |
| Patient name: | | | Pa | tient ID #: | | | DOB: |
| Street address: | | | | | | | |
| Apt #: | City/state/zip: | | | Ph | ione: | | |
| PRESCRIBER INFO | DRMATION | | | | | | |
| Prescriber name: | | | | | | | |
| Specialty: | | | | NPI: | | | State license #: |
| Street address: | | | | | | | |
| Suite #: | City/state/zip: | | | | | | |
| Phone: | | | Fa | X: | | | |
| CLINICAL INFORM | 1ATION | | | | | | |
| Medication requested: | | | No | on-Preferred N | lodications: | | |
| Preferred Medications: Actemra (tocilizumab) Syringe | | ren ryyringe ge ettor et qyh) | □ Actemra (tocilizumab) Actpen □ Adalimumab-adaz(CF) 100 mg/ml Pen □ Adalimumab-adaz(CF) 100 mg/ml Syringe □ Amjevita(CF) (adalimumab-atto) 50 mg/ml Autoinjector □ Amjevita(CF) (adalimumab-atto) 50 mg/ml Syringe □ Arcalyst (rilonacept) Vial □ Cimzia (certolizumab pegol) Syringe □ Cosentyx (secukinumab) Pen □ Cosentyx (secukinumab) Syringe □ Cyltezo(CF) (adalimumab-adbm) 50 mg/ml Pen □ Cyltezo(CF) (adalimumab-adbm) 50 mg/ml Syringe □ Entyvio (vedolizumab) Vial □ Hulio(CF) (adalimumab-fkjp) 50 mg/ml Pen □ Hulio(CF) (adalimumab-fkjp) 50 mg/ml Syringe □ Hyrimoz(CF) (adalimumab-adaz) 100 mg/ml Syringe □ Hyrimoz(CF) (adalimumab-adaz) 100 mg/ml Syringe □ Idacio(CF) (adalimumab-aacf) 50 mg/ml Pen □ Idacio(CF) (adalimumab-aacf) 50 mg/ml Syringe □ Idacio(CF) (adalimumab-aacf) 50 mg/ml Syringe □ Idacio(CF) (adalimumab-aacf) | | 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) Syringe □ Styrizi (risankizumab) Vial □ Sotyktu (deucravacitinib) Tablet □ Spevigo (spesolimab-sbzo) Vial □ Stelara (ustekinumab) Syringe □ Stelara (ustekinumab) Syringe □ Stelara (ustekinumab) Autoinjector □ Tremfya (guselkumab) Autoinjector □ Tremfya (folacitinib) Solution □ Yuflyma(CF) (adalimumab-aaty) 100 □ mg/ml Autoinjector □ Yuflyma(CF) (adalimumab-aaty) 100 □ mg/ml Syringe | |
| STARTER PACK requested | (strength/formulatio | nn): | MA | AINTENANCE pr | oduct/packaging reque | ested (str | ength/formulation): |
| Quantity per fill: | | Refills: | Qu | antity per fill: | | R | defills: |
| Directions: | | | Dir | Directions: | | | |
| Diagnosis (submit documentation): | | | | Dx code (required): | | В | leneficiary weight: |
| Is the beneficiary currently being treated with the requested medication? | | | | ☐ Yes – 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)? | | | | ☐ Yes ☐ No If prescriber is not a specialist, submit documentation of consultation. | | | |



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|---|----------------------|------------------------------------|--|--|--|--|--|
| PHARMACY INFORMATION (Prescriber to identify the pharmacy that is to dispense the medication): | | | | | | | |
| Deliver to: ☐ Patient's Home | ☐ Physician's Office | ☐ Patient's Preferred Pharmacy Nar | ne: | | | | |
| NPI#: | | | | | | | |
| Pharmacy Phone #: | | | Pharmacy Fax #: | | | | |
| ☐ I acknowledge that the patient agrees with the pharmacy chosen for delivery of this medication. | | | | | | | |

| ∃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 submit documentation for each item. |
| INI | 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 |
| Diag | nosis |
| | ALL diagnoses: □ 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 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, 6-MP, MTX) |
| 7. | Familial Mediterranean fever: ☐ Tried and failed a 3-month trial of or has a contraindication or an intolerance to colchicine at maximally tolerated doses |
| 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 corticosteroids Has a medical reason why repeated courses of corticosteroids are not appropriate |
| 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) |



CYTOKINE AND CAM ANTAGONISTS PRIOR AUTHORIZATION FORM **INITIAL REQUESTS (continued)** 11. Juvenile idiopathic arthritis: ☐ Has systemic disease with active systemic features \square Has disease associated with any of the following: ☐ Positive anti-CCP antibodies □ Positive rheumatoid factor \square 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 12. 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) ☐ 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 ☐ Has severe disease ☐ Has comorbid moderate-to-severe nail psoriasis ☐ Has comorbid active inflammatory bowel disease ☐ Tried and failed a 3-month trial of or has a contraindication or an intolerance to conventional DMARDs (e.g., AZA, leflunomide, MTX, etc.) ☐ 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) 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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