



## envolve pa health wellness. Prior Authorization Request Form for Cytokine and CAM **Prior Authorization Request Antagonists**

## FAX this completed form to (877) 386-4695

IL MEMBER INFORMATION	Man requests to: Envolve r narmacy solution	пз і А Бера						
Prescriber Specialty:   Identification #:   Office Contact Name:   Group #:   Orton Name:   Date of Birth:	I. PROVIDER INFORMATION		II. M	II. MEMBER INFORMATION				
Office Contact Name:   Group #: Group Name:   Date of Birth:   Phone #:   Medication Allergies:   Phone #:   Date of Birth:	Prescriber Name:		Memb	Member Name:				
Group Name:   Fax #:   Medication Allergies:   Qty. per Day:   W. REQUIRED DOCUMENTION (Detailed medical record documentation demonstrating evidence for each item must be submitted with prior authorization request)  Specify diagnosis & diagnosis code relevant to this request:   Dx/Dx Code:	Prescriber Specialty:		Identification #:					
Phone #:	Office Contact Name:		Group #:					
Phone #:    III. DRUG INFORMATION (One drug request per form)	Group Name:		Date of Birth:					
Drug name and strength:    Dosage Interval (sig):   Qty. per Day:	Fax #:		Medic	Medication Allergies:				
Dosage Interval (sig):   Qty. per Day:	Phone #:							
IV. REQUIRED DOCUMENTION (Detailed medical record documentation demonstrating evidence for each item must be submitted with prior authorization request)  Specify diagnosis & diagnosis code relevant to this request:  Does the member have any contraindications to the prescribed medication?  All potential drug interactions have been addressed by the prescriber such as discontinuation or dose reduction of interacting medication or counseling the member about the risks associated with the use of both interacting medications.  Requests for all non-preferred medications: Does the member have a history of trial and failure of or contraindication or intolerance to the preferred Cytokine and CAM Antagonist? Refer to https://www.dhs.pa.gov/providers/Pharmacy-Services/Pages/Quantity-Limits-and-Daily-Dose-Limits.aspx), please provide supporting information:  SUBMIT MEDICAL RECORD INFORMATION FOR EACH APPLICABLE ITEM.  If not prescribed by one of the following specialist, gastroenterologist, dermatologist, rheumatologist, ophthalmologist, immunologist, genetic specialist, etc., please indicate a specialist consulted:  The requested medication will NOT be use concurrently with another Cytokine and CAM Antagonist  For Cytokine and CAM Antagonist associated with an increased risk of infection according to FDA-approved package labeling:  Member is up to date with immunizations in accordance with CDC Advisory Committee on Immunization Practices (ACIP) OR  Has a plan for receiving CDC ACIP recommended immunizations  Member has and hepatitis B screening (sAb, sAg and cAb)  If screening indicates a risk of hepatitis B trivus reactivation, a follow-up plan to address this risk OR	III. DRUG INFORMATION (One drug reque	st per forn	n)					
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☐ If screening indicates a risk of hepatitis B virus reactivation, a follow-up plan to address this risk OR								

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	package labeling:
	<ul> <li>Member was evaluated for a history of prior suicide attempt, bipolar disorder, or major depressive disorder</li> <li>Member will be monitored for behavioral and mood changes as recommended in the FDA approved package</li> </ul>
	labeling
	IT MEDICAL RECORD INFORMATION FOR EACH APPLICABLE ITEM.
	'-ONSET STILL'S DISEASE:
Ш	Member has predominantly systemic disease:
	Member has steroid-dependent disease and will be using requested medication with the intent of discontinuation or
	decreasing the dose of systemic steroids
	Member has tried and failed or has a contraindication or intolerance to corticosteroids (medication, start date and end date):
	Member has predominantly joint disease:
	☐ Member has tried and failed or has a contraindication or intolerance to conventional non-biologic DMARDs (eg,
	Methotrexate)(medication, start date and end date):
_	OSING SPONDYLITIS & NON-RADIOGRAPHIC AXIAL SPONDYLOARTHRITIS:
BEHCE	(medication, start date and end date):
	Member has recurrent oral ulcers associated with Behcet's syndrome
	Member has tried and failed or has a contraindication or intolerance to topical corticosteroids (medication, start date
	and end date):
	Member has tried and failed at least 3 month trial or has a contraindication or intolerance to colchicine (medication,
	start date and end date):
	NIC PSORIASIS:
	Member has moderate to severe chronic psoriasis
	Member has involvement of critical areas of the body (eg, skin folds, face, genitals)
╽	Member has psoriasis causing significant disability or impaired physical or mental functioning
	Member has tried and failed 3 month trial or has a contraindication or intolerance to oral systemic therapy (eg,
	Azathioprine, Cyclosporine, Acitretin) (medication, start date and end date):
	Member has tried and failed or has a contraindication or intolerance to topical corticosteroids (medication, start date
	and end date):
	Member has tried and failed or has a contraindication or intolerance to ultraviolet light therapy (start date and end
CDOIII	date):
	J'S DISEASE:  Mamban bas madanata ta sayana Chabn's diseasa
	Member has moderate to severe Crohn's disease Member has disease associated with high-risk or poor prognostic features
	Member has tried and failed to achieve remission with or has a contraindication or intolerance to an induction course of
	corticosteroids (medication, start date and end date):
	Member has tried and failed to maintain remission with or has a contraindication or intolerance to an
_	immunomodulators (Azathioprine, Mercaptopurine, Methotrexate) (medication, start date and end
	date):
	remission
FAMIL	IAR MEDITERRANEAN FEVER:
	Member has tried and failed at least 3 month trial or has a contraindication or intolerance to colchicine (medication, start date and end date):
GAINT	CELL ARTERITIS:
	Member has tried and failed or has a contraindication or intolerance to systemic, corticosteroids (medication, start date
	and end date):
	·
	Member has glucocorticoid-dependent disease and will be using requested medication with the intent of discontinuing
	or decreasing the dose of steroids
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HIDRA	DENITIS SUPPURATIVA:
	Member has Hurley stage II or III disease
	Member has tried and failed 3 month trial or has a contraindication or intolerance to topical Clindamycin (medication, start date and end date):
	Member has tried and failed or has a contraindication or intolerance to systemic antibiotic (eg. doxycycline, minocycline, tetracycline, clindamycin) (medication, start date and end date):
IUVEN	ILE IDIOPATHIC ARTHRITIS (JIA):
	Member has systemic juvenile idiopathic arthritis with active systemic features
	Member has a diagnosis of juvenile idiopathic arthritis associated with a high disease activity or one or more poor prognostic feature(s)
	Member has active sacrooilitis and/or enthesitis:
	☐ Member has tried and failed a 2 week trial or has a contraindication or intolerance to oral NSAIDs (medication, start date and end date):
<b>PSORI</b>	ATIC ARTHRITIS:
	Member has severe active psoriatic arthritis
	Member has concomitant moderate to severe nail disease
	Member has predominantly axial disease and/or enthesitis:
	Member has tried and failed a 2 week trial or has a contraindication or intolerance to 2 different oral NSAIDs (medication, start date and end date):
	Member has predominantly peripheral disease:
	☐ Member has tried and failed a 8 week trial or has a contraindication or intolerance to conventional non-biologic DMARDs (Azathioprine, Leflunomide, Methotrexate, Sulfasalazine)(medication, start date and end date):
RHEUN	MATOID ARTHRITIS:
	Member has moderate to severe active rheumatoid arthritis
=	Member has tried and failed a 3 month trial or has a contraindication or intolerance to conventional DMARDs (Azathioprine, Leflunomide, Methotrexate, etc) (medication, start date and end date):
	ATIVE COLITIS (UC):
	Member has moderate to severe ulcerative colitis
	Member has mild disease associated with high-risk or poor prognostic features
	Member has tried and failed to achieve remission with or has a contraindication or intolerance to an induction course of corticosteroids (medication, start date and end date):
	Member has tried and failed to maintain remission with or has a contraindication or intolerance to an immunomodulators (Azathioprine, Cyclosporine, Mercaptopurine, Methotrexate) (medication, start date and end date):
	Member has achieved remission with the requested medication and will be using as maintenance therapy to maintain remission
IIVEIT	IS (NON-INFECTIOUS):
	Member has comorbid juvenile idiopathic arthritis
	•
	Member has steroid-dependent disease
	Member has tried and failed or has a contraindication or intolerance to systemic, topical, intraocular or periocular corticosteroids (medication, start date and end date):
	Member has tried and failed or has a contraindication or intolerance to systemic immunosuppressive (Azathioprine, Methotrexate, Mycophenolate Mofetil, etc) (medication, start date and end date):
	Member has corticosteroid-dependent uveitis and will be using requested medication with the intent of discontinuing
	or decreasing the dose of steroids
	YST (RILONACEPT):  Mambar has tried and failed to maintain remission with or has a centraindication or intelevance to Vinerat (analyzing).
_	Member has tried and failed to maintain remission with or has a contraindication or intolerance to Kineret (anakinra) (medication, start date and end date):
	Member has taken the requested non-preferred antipsychotic in the past 90 days:

ILARIS (CANAKINUMAB):								
Member has tried and failed to maintain remission with or has a contraindication or intolerance to Kineret (anakinra) (medication, start date and end date):								
☐ Member has taken the requested non-preferred antipsychotic in the past 90 days:								
INFL <u>ix</u> imab product other than avsola (inflixima								
Member has tried and failed to maintain remission with or has a contraindication or intolerance to Avsola (infliximab-axxq) (medication, start date and end date):								
☐ Member has taken the requested non-preferred antip	sychotic in the past 90 days:							
RENEWAL REQUEST:								
Member has experienced improvement in disease activity and/or level of functioning since initiating therapy with requested medication:								
IV. ADDITIONAL RATIONALE FOR REQUEST / PERT								
Appropriate clinical information to augment the received or	Duaridan Cianatura.	Data						
Appropriate clinical information to support the request on the basis of medical necessity must be submitted.	Provider Signature:	Date:						
the basis of medical hecessity must be submitted.								

Envolve Pharmacy Solutions will respond via fax or phone within 24 hours.

Requests for prior authorization (PA) requests must include member name, ID#, and drug name. Please include lab reports with requests when appropriate (e.g., Culture and Sensitivity; Hemoglobin A1C; Serum Creatinine; CD4; Hematocrit; WBC, etc.)