AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete</u>, correct, or legible, the authorization process can be delayed.

Drug Requested: Kineret® (anakinra)

MEMBER & PRESCRIBER INI	FORMATION: Authorization may be delayed if incomplete.
Member Name:	
Member AvMed#:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	Date:
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Author	rization may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code:
Weight:	Date:
mmunomodulator (e.g., Dupixent, Entyvi	se of concomitant therapy with more than one biologic io, Humira, Rinvoq, Stelara) prescribed for the same or different gational. Safety and efficacy of these combinations has NOT been
	elow all that apply. All criteria must be met for approval. To ation, including lab results, diagnostics, and/or chart notes, must be
□ Diagnosis: Moderate-to-Seven Dosing: SubQ: 100 mg daily	re Active Rheumatoid Arthritis
Authorization Criteria: (Length	of authorization is indefinite for this indication only)
☐ Member has a diagnosis of modera	ate-to-severe active rheumatoid arthritis
☐ Prescribed by or in consultation w	rith a Rheumatologist

				led at least <u>ONE</u> of the following DMARD therapies for at least rt notes or pharmacy paid claims)	ast three (3)
		hyd	roxychloroquine		
		leflu	unomide		
		met	hotrexate		
		sulf	asalazine		
	Me	embe	r meets ONE of t	he following:	
		Mei	mber tried and fai	led, has a contraindication, or intolerance to <u>TWO</u> of the <u>PRE</u> fied by chart notes or pharmacy paid claims):	FERRED
			Actemra® SC	☐ adalimumab product: Humira [®] , Cyltezo [®] or Hyrimoz [®]	□ Enbrel®
			Rinvoq®	□ Xeljanz [®] /XR [®]	
		Abb		C's starting with 83457 are not approved, NDC's starting with Hyrimoz NDC's starting with 83457 are not approved, NDC at a preferred	`
		indi	cates <u>at least a 90</u>	ablished on Kineret [®] for at least 90 days <u>AND</u> prescription clands and the past 130 macy paid claims)	•
	_		is: Systemic JubQ: 100 mg dail	uvenile Idiopathic Arthritis (SJIA) ly	
Init	ial A	Auth	norization: 12	months	
	Da	te of	diagnosis must be	e noted:	
				rial and failure of NSAIDs and corticosteroids for > 3 months (verified by chart notes or pharmacy paid claims)	consecutively
				2 active joints with concomitant fever for at least 5 days and dosed at 0.5 mg/kg/day or 30 mg/day within the last 3 months	
		embe luest		ever > 38° C or 100.4° F for at least 2 weeks within the last 2 to	months of this
	Me	embe	r must have ONE	of the following measurements of active disease:	
		Mei	mber must have h	ad CRP (>15 mg/L) within the last 2 months of this request	
		Mei	mber must have ha	ad ESR (>45 mm/hr) within the last 2 months of this request	
	_		is: Adult-onse ubQ: 100 mg dail	t Still's disease (AOSD)	
Initial Authorization: 12 months					
	Me	embe	r must be at least	18 years of age	

	Member must meet TWO of the following:
	☐ Fever >39°C, lasting 1 week or longer
	☐ Arthralgia or arthritis, lasting 2 weeks or longer
	□ Typical rash
	☐ Leukocytosis >10,000/mm³ with >80% polymorphonuclear cells
	Disease activity based on DAS28 of \geq 3.2 at screening
	Member must have ONE of the following measurements of active disease:
	☐ Member must have had CRP (>15 mg/L) within the last 2 months of this request
	☐ Member must have had ESR (>45mm/hour) within the last 2 months of this request
	Member must have had ≥ 2 joints that are painful/swollen for at least 2 weeks within the last 3 months of this request
	Member must have had trial and failure with at a least 1 week of glucocorticoids (dose: ≤ 10 mg/day prednisolone equivalent) <u>AND</u> at least 4 weeks of NSAIDs within the last 3 months of this request
□ D	Diagnosis: Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS)
Initi	al Authorization: 12 months
	Prescribed by or in consultation with a Rheumatologist or Immunologist with expertise in the diagnosis of TRAPS
	Member has a diagnosis of TRAPS with genetic confirmation of the TNFRSF1A gene mutation
	Member has had chronic or recurrent disease resulting in six (6) flares within a 12-month time frame (submit chart notes)
	Provider must submit labs documenting the member's CRP level >10 mg/L which is indicative of active disease (submit labs collected within the last 30 days)
	Member must have trial and failure of NSAIDs and corticosteroids within the last 6 months (verified by chart notes or pharmacy paid claims)
	Diagnosis: Hyperimmunoglobulin D Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD)
<u>Initi</u>	al Authorization: 12 months
	Prescribed by or in consultation with a Rheumatologist or Immunologist with expertise in the diagnosis of HIDS/MKD
	Provider must submit genetic confirmation of HIDS (i.e. DNA analysis or enzymatic studies showing mutations in the MVK gene or markedly reduced mevalonate kinase activity)
	Member must have a history of \geq three (3) febrile acute flares within a 6-month period when not receiving prophylactic treatment
	Provider must submit labs documenting the member's CRP level >10mg/L which is indicative of active disease (submit labs collected within the last 30 days)

I	Diagnosis: Familial Mediterranean Fever (FMF) Maximum Dosing: SubQ: 100 mg daily Children ≥ 2 years and Adolescents: SubQ: 2 mg/kg/dose once daily
Init	tial Authorization: 12 months
	Prescribed by or in consultation with a Rheumatologist or Immunologist with expertise in the diagnosis of FMF
	Member must have Type 1 disease characterized by recurrent and short episodes of inflammation and serositis with an average of at least one documented acute FMF attack per month during the previous 6 months and lasting approximately 12 to 72 hours
	Provider must submit genetic confirmation of active Type 1 FMF disease (i.e., MEFV gene exon 10 mutation)
	Provider must submit labs documenting the member's CRP level >10mg/L which is indicative of active disease (submit labs collected within the last 30 days)
	Member must have trial and failure of maximally dosed colchicine (children-2 mg/day or adults-3 mg/day)
i i	Diagnosis: Cryopyrin-associated periodic syndromes (CAPS) Dosing: Children, and Adolescents: SubQ: Initial: 1 to 2 mg/kg/day in 1 to 2 divided doses; adjust dose n 0.5 to 1 mg/kg increments as needed to control inflammation; usual maintenance dose: 3 to 4 mg/kg/day; maximum daily dose: 8 mg/kg/day tial Authorization: 12 months
	Prescribed by or in consultation with a Rheumatologist or Immunologist with expertise in the diagnosis of CAPS
	Member must have at least <u>TWO</u> of any of the CAPS-typical symptoms: □ urticaria-like rash □ cold-triggered episodes □ sensorineural hearing loss □ musculoskeletal symptoms □ chronic aseptic meningitis □ skeletal abnormalities
	Member has elevated serum levels which are indicative of active disease: (submit labs collected within the last 30 days)
	□ C-Reactive Protein (CRP): AND □ Serum Amyloid A (SAA):
	Member has documented laboratory evidence of a genetic mutation in the Cold-Induced Auto-inflammatory Syndrome 1 (CIAS1), also known as NLRP3 (R26OW, T348M, D303N, E311K, M662T, A439V, D305N, T436N, T4361) (submit genetic testing results)

 Member has a diagnosis of <u>ONE</u> of the following: Familial Cold Auto-inflammatory Syndrome (FCAS) Muckle- Wells Syndrome (MWS) Neonatal-Onset Multisystem Inflammatory Disease (NOMID)
□ Diagnosis: Deficiency of interleukin 1 receptor antagonist (DIRA) Dosing: Children, and Adolescents: SubQ: Initial: 1 to 2 mg/kg/dose once daily; may titrate in 0.5 to 1 mg/kg increments up to a maximum dose of 8 mg/kg/dose
Initial Authorization: 12 months
☐ Prescribed by or in consultation with a Rheumatologist or Immunologist with expertise in the diagnosis of DIRA
☐ Member is <u>NOT</u> receiving another IL1 antagonist medication (e.g., Ilaris or Araclyst)
☐ Member has <u>ONE</u> of the following: pustular dermatitis, osteomyelitis, vertebral destruction (submit chart note documentation)
☐ Member has elevated serum levels indicative of active disease (submit labs collected within the last 3 days)
□ C-Reactive Protein (CRP): <u>OR</u> □ Erythrocyte Sedimentation Rate (ESR):
Reauthorization: 12 months. Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied. Note: Reauthorization criteria is applicable for all diagnoses EXCEPT Rheumatoid Arthritis.
☐ Member has experienced an absence of unacceptable toxicity from the drug [e.g., hypersensitivity reactions, serious infections (include but not limited to tuberculosis), and macrophage activation syndrome (MAS)]
☐ Member is receiving ongoing monitoring for presence of TB or other active infections
☐ Member has experienced disease response as indicated by improvement in member's symptoms from baseline AND improvement of CRP and SAA serum levels (both levels are <10 mg/L) (submit labs collected within the last 30 days)
Medication being provided by Specialty Pharmacy – Proprium Rx

Use of samples to initiate therapy does not meet step edit/preauthorization criteria.

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.