AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) 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 (including phone and fax #s) on this form is correct. <u>If the information provided is not</u> complete, correct, or legible, the authorization process can be delayed.

Drug Requested: Kineret[®] (anakinra)

MEMBER & PRESCRIBER INFORMATION: 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:	
NPI #:		
DRUG INFORMATION: Authorizatio	n may be delayed if incomplete.	
Drug Name/Form/Strength:		
	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight (if applicable):	Date weight obtained:	
immunomodulator (e.g., Dupixent, Entyvio, Hu	concomitant therapy with more than one biologic umira, Rinvoq, Stelara) prescribed for the same or different nal. Safety and efficacy of these combinations has <u>NOT</u> been	
Will the member be discontinuing a previously prescribed biologic if approved for requested medication?		
• If yes, please list the medication that will be approval along with the corresponding effe	e discontinued and the medication that will be initiated upon active date.	
Medication to be discontinued:	Effective date:	
	Effective date:	

provided or request may be denied.

Diagnosis: Moderate-to-Severe Active Rheumatoid Arthritis Dosing: SubQ: 100 mg daily

Authorization Criteria: (Length of authorization is indefinite for this indication only)

- D Member has a diagnosis of moderate-to-severe active **rheumatoid arthritis**
- **D** Prescribed by or in consultation with a **Rheumatologist**
- □ Member has tried and failed at least <u>ONE</u> of the following **DMARD** therapies for at least three <u>(3)</u> <u>months</u> (verified by chart notes or pharmacy paid claims):
 - □ hydroxychloroquine
 - □ leflunomide
 - □ methotrexate
 - □ sulfasalazine
- □ Member meets <u>ONE</u> of the following:
 - □ Member tried and failed, has a contraindication, or intolerance to <u>**TWO**</u> of the <u>**PREFERRED**</u> biologics below (verified by chart notes or pharmacy paid claims):

Preferred adalimumab product*	□ Enbrel [®]
□ Rinvoq [®] /Rinvoq [®] LQ	 Preferred tocilizumab product: Actemra[®] SC or Tyenne[®] SC
$\Box \text{Xeljanz}^{\circledast}/\text{XR}^{\circledast}$	

*<u>NOTE</u>: COMM/FAMIS preferreds = Humira/Cyltezo/Yuflyma - Humira NDC's starting with 83457 are not approved, NDC's starting with 00074 (MFG: Abbvie) are preferred; SG/IP/HIX preferreds = Simlandi or adalimumab-adbm

Member has been established on Kineret[®] for at least 90 days <u>AND</u> prescription claims history indicates <u>at least a 90-day supply of Kineret was dispensed within the past 130 days</u> (verified by chart notes or pharmacy paid claims)

 Diagnosis: Systemic Juvenile Idiopathic Arthritis (SJIA) Dosing: SubQ: 100 mg daily

Initial Authorization: 12 months

- Date of diagnosis must be noted:
- □ Member must have had trial and failure of NSAIDs and corticosteroids for > 3 months consecutively within the last 4 months (verified by chart notes or pharmacy paid claims)
- □ Member must have had ≥ 2 active joints with concomitant fever for at least 5 days and trial of prednisone or equivalent dosed at 0.5 mg/kg/day or 30 mg/day within the last 3 months of this request
- □ Member must have had fever > 38° C or 100.4° F for at least 2 weeks within the last 2 months of this request

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- □ Member must have <u>ONE</u> of the following measurements of active disease:
 - \Box Member must have had CRP (>15 mg/L) within the last 2 months of this request
 - □ Member must have had ESR (>45 mm/hr) within the last 2 months of this request

Diagnosis: Adult-onset Still's disease (AOSD) Dosing: SubQ: 100 mg daily

Initial Authorization: 12 months

- □ Member must be at least 18 years of age
- □ Member must meet <u>**TWO**</u> of the following:
 - \Box Fever >39°C, lasting 1 week or longer
 - □ Arthralgia or arthritis, lasting 2 weeks or longer
 - □ Typical rash
 - \Box Leukocytosis >10,000/mm³ with >80% polymorphonuclear cells
- □ Disease activity based on DAS28 of \geq 3.2 at screening
- □ Member must have <u>ONE</u> 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: $\leq 10 \text{ mg/day}$ prednisolone equivalent) <u>AND</u> at least 4 weeks of NSAIDs within the last 3 months of this request

Diagnosis: Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS) <u>Initial Authorization</u>: 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)

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Diagnosis: Hyperimmunoglobulin D Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD)

Initial 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)
- □ Diagnosis: Familial Mediterranean Fever (FMF) Maximum Dosing: SubQ: 100 mg daily Children ≥ 2 years and Adolescents: SubQ: 2 mg/kg/dose once daily

Initial 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 >10 mg/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)

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 in 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

Initial 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 Autoinflammatory 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 30 days)
 - □ C-Reactive Protein (CRP): ____ □ Erythrocyte Sedimentation Rate (ESR): _

<u>Reauthorization</u>: 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. <u>Note</u>: 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 <u>AND</u> improvement of CRP and SAA serum levels (both levels are <10 mg/L) (submit labs collected within the last 30 days)</p>

Medication being provided by Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required. **Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.** *<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>*