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, correct, or legible, the authorization process can be delayed.</u>

□ Rinvoq [®] (upadacitinib)	□ Rinvoq® LQ (upadacitinib)		
MEMBER & PRESCRIBER INFO	RMATION: Authorization may be delayed if incomplete.		
Member Name:			
Member AvMed #:	er AvMed #: Date of Birth:		
Prescriber Name:			
Prescriber Signature:			
Office Contact Name:			
Phone Number:			
NPI #:			
DRUG INFORMATION: Authorization	on 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, H	f concomitant therapy with more than one biologic fumira, Rinvoq, Stelara) prescribed for the same or different onal. Safety and efficacy of these combinations has NOT been		
Will the member be discontinuing a previously prescribed biologic if approved for requested medication?			
	□ Yes OR □ No		
• If yes, please list the medication that will be approval along with the corresponding effective.	be discontinued and the medication that will be initiated upon ective date.		
Medication to be discontinued:	Effective date:		
Medication to be initiated:	Effective date:		

CLINICAL CRITERIA: 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.

□ Diagnosis: Moderate-to-Severe Rheumatoid Arthritis Dosing: Oral: Rinvoq® 15 mg once daily			
☐ Member has a diagnosis of moderate-to-severe rheumatoid arthritis			
☐ Prescribed by or in consultation with a Rheumatologist			
☐ Member is 18 years of age or older			
☐ Member has tried and failed at least <u>ONE</u> of the following DMARD therapies for at least <u>three (3)</u> <u>months</u>			
□ hydroxychloroquine			
□ methotrexate			
□ sulfasalazine			
☐ Member meets <u>ONE</u> of the following:			
☐ Member tried and failed, has a contraindication, or intolerance to ONE of the following:			
□ <u>ONE</u> preferred adalimumab product [<u>NOTE</u> : <u>COMM/FAMIS</u> preferreds = <u>Humira/Cyltezo/Yuflyma</u> - Humira NDC's starting with 83457 are not approved, NDC's starting with 00074 (MFG: Abbvie) are preferred; <u>SG/IP/HIX</u> preferreds = <u>Simlandi</u> or adalimumab-adbm			
□ Enbrel [®]			
Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Moderate-to-Severe Rheumatoid Arthritis:			
Member has been established on Rinvoq [®] for at least 90 days <u>AND</u> prescription claims history indicates <u>at least a 90-day supply of Rinvoq was dispensed within the past 130 days</u> (verified least notes or pharmacy paid claims)			
☐ Member is <u>NOT</u> receiving Rinvoq [®] in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants			
□ Diagnosis: Active Psoriatic Arthritis			

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Dosing: Oral: Rinvoq® or Rinvoq® LQ			
Patient Age	Patient Weight	Rinvoq® LQ	Rinvoq®
	10 kg to < 20 kg	3 mg (3 mL) twice daily	Not Recommended
2 to < 18 years of age	20 kg to < 30 kg	4 mg (4 mL) twice daily	Not Recommended
	> 30 kg	6 mg (6 mL) twice daily	15 mg once daily
≥ 18 years of age	N/A	6 mg (6 mL) twice daily	15 mg once daily

Member has a diagnosis of active psoriatic arthritis		
Prescribed by or in consultation with a Rheumatologist		
Member is 2 years of age or older		
Member has tried and failed at least <u>ONE</u> of the following DMARD therapies for at least <u>three</u> (<u>months</u>		
□ cyclosporine		
□ leflunomide		
□ methotrexate		
□ sulfasalazine		
Member meets ONE of the following:		
☐ Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following:		
□ <u>ONE</u> preferred adalimumab product [<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]		
□ Enbrel [®]		
 Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Active Psoriatic Arthritis: 		
Member has been established on Rinvoq® for at least 90 days <u>AND</u> prescription claims history indicates <u>at least a 90-day supply of Rinvoq was dispensed within the past 130 days</u> (verified by chart notes or pharmacy paid claims)		
Member is <u>NOT</u> receiving Rinvoq [®] in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants		
viagnosis: Moderate-to-Severe Atopic Dermatitis osing: Oral: Rinvoq® 15 mg once daily; may increase to 30 mg once daily if inadequate response		
Member has a diagnosis of <u>moderate to severe atopic dermatitis</u> with disease activity confirmed by <u>ONE</u> of the following (chart notes documenting disease severity and BSA involvement must be included):		
□ Body Surface Area (BSA) involvement >10%		
☐ Eczema Area and Severity Index (EASI) score ≥ 16		
□ Investigator's Global Assessment (IGA) score ≥ 3		
☐ Scoring Atopic Dermatitis (SCORAD) score ≥ 25		
Prescribed by or in consultation with an Allergist, Dermatologist or Immunologist		
Member is 12 years of age or older		
Member weighs at least 40 kg		
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	Member is NOT receiving Rinvoq [®] in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants			
	Member has tried and failed at least <u>ONE</u> of the following DMARD therapies for at least <u>three (3)</u> months			
		aza	athioprine	
		cyc	closporine	
		me	ethotrexate	
		my	cophenolate mofetil	
☐ Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following topical the (chart notes documenting contraindication(s) or intolerance must be attached; trials will be verified using pharmacy claims and/or submitted chart notes):				
			days (14 days for very high potency) of therapy with <u>ONE</u> medium to very-high potency topical rticosteroid in the past 180 days	
			days of therapy with <u>ONE</u> of the following topical calcineurin inhibitors in the past 180 days: tacrolimus 0.03 % or 0.1% ointment	
		_ _	pimecrolimus 1% cream (requires prior authorization)	
d D	osag isco	ge o	Oral: Rinvoq [®] : Induction - 45 mg once daily for 8 weeks; Maintenance - 15 mg once daily. A f 30 mg once daily may be considered for patients with refractory, severe or extensive disease. The interpretation of the second patients with the 30 mg dose. Use the lowest effective dose of maintain response.	
	Me	emb	er has a diagnosis of moderate-to-severe ulcerative colitis	
	Pre	escr	ibed by or in consultation with a Gastroenterologist	
	Me	emb	er is 18 years of age or older	
	Member meets <u>ONE</u> of the following:			
			ember has tried and failed budesonide or high dose steroids (40-60 mg prednisone)	
			ember has tried and failed at least ONE of the following DMARD therapies for at least three (3) onths	
			5-aminosalicylates (balsalazide, olsalazine, sulfasalazine)	
			oral mesalamine (Apriso, Asacol/HD, Delzicol, Lialda, Pentasa)	
	Me	emb	er meets ONE of the following:	
			ember tried and failed, has a contraindication, or intolerance to ONE of the following:	
			ONE preferred adalimumab product [NOTE: 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]	
			Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Moderate-to-Severe Ulcerative Colitis:	

_	 Member has been established on Rinvoq® for at least 90 days AND prescription claims history indicates at least a 90-day supply of Rinvoq was dispensed within the past 130 days (verified by chart notes or pharmacy paid claims) Member is NOT receiving Rinvoq® in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants
E A d	Diagnosis: Moderate-to-Severe Active Crohn's Disease (CD) Dosing: Oral: Rinvoq®: Induction - 45 mg once daily for 12 weeks; Maintenance -15 mg once daily. I dosage of 30 mg once daily may be considered for patients with refractory, severe or extensive isease. Discontinue if an adequate response is not achieved with the 30 mg dose. Use the lowest effective dose needed to maintain response.
	Member has a diagnosis of moderate-to-severe Crohn's disease
	Prescribed by or in consultation with a Gastroenterologist
	Member is 18 years of age or older
	Member meets ONE of the following:
	☐ Member has tried and failed budesonide or high dose steroids (40-60 mg prednisone)
	☐ Member has tried and failed at least <u>ONE</u> of the following DMARD therapies for at least <u>three (3)</u> months
	□ 5-aminosalicylates (balsalazide, olsalazine, sulfasalazine)
	oral mesalamine (Apriso, Asacol/HD, Delzicol, Lialda, Pentasa)
	Member meets ONE of the following:
	☐ Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following:
	□ <u>ONE</u> preferred adalimumab product [<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]
	☐ Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Moderate-to-Severe Active Crohn's Disease:
	Member has been established on Rinvoq [®] for at least 90 days <u>AND</u> prescription claims history indicates <u>at least a 90-day supply of Rinvoq was dispe</u>
	Member is <u>NOT</u> receiving Rinvoq [®] in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants
	Diagnosis: Active Ankylosing Spondylitis Dosing: Oral: Rinvoq® 15 mg once daily
	Member has a diagnosis of active ankylosing spondylitis
	Prescribed by or in consultation with a Rheumatologist

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	Member is 18 years of age or older
	Member tried and failed, has a contraindication, or intolerance to TWO NSAIDs
	Member meets ONE of the following:
	☐ Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following:
	ONE preferred adalimumab product [NOTE: 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]
	□ Enbrel [®]
	 Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Active Ankylosing Spondylitis:
	Member has been established on Rinvoq® for at least 90 days <u>AND</u> prescription claims history indicates <u>at least a 90-day supply of Rinvoq was dispensed within the past 130 days</u> (verified by chart notes or pharmacy paid claims)
	Member is <u>NOT</u> receiving Rinvoq [®] in combination with other JAK inhibitors, biologic immunomodulators, or with other immunosuppressants
	Diagnosis: Active Non-Radiographic Axial Spondyloarthritis Dosing: Oral: Rinvoq® 15 mg once daily
	Member has a diagnosis of active non-radiographic axial spondyloarthritis
	Prescribed by or in consultation with a Rheumatologist
	Member is 18 years of age or older
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	Member has at least ONE of the following objective signs of inflammation:
	Member has at least <u>ONE</u> of the following objective signs of inflammation: □ C-reactive protein [CRP] levels above the upper limit of normal
_	 □ C-reactive protein [CRP] levels above the upper limit of normal □ Sacroiliitis on magnetic resonance imaging [MRI] (indicative of inflammatory disease, but without
_ _	 C-reactive protein [CRP] levels above the upper limit of normal Sacroiliitis on magnetic resonance imaging [MRI] (indicative of inflammatory disease, but without definitive radiographic evidence of structural damage on sacroiliac joints)
	 C-reactive protein [CRP] levels above the upper limit of normal Sacroiliitis on magnetic resonance imaging [MRI] (indicative of inflammatory disease, but without definitive radiographic evidence of structural damage on sacroiliac joints) Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> NSAIDs
	 C-reactive protein [CRP] levels above the upper limit of normal Sacroilitis on magnetic resonance imaging [MRI] (indicative of inflammatory disease, but without definitive radiographic evidence of structural damage on sacroiliac joints) Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> NSAIDs Member meets <u>ONE</u> of the following: Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following: Cimzia[®] Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Active Non-Radiographic Axial Spondyloarthritis:
	 C-reactive protein [CRP] levels above the upper limit of normal Sacroiliitis on magnetic resonance imaging [MRI] (indicative of inflammatory disease, but without definitive radiographic evidence of structural damage on sacroiliac joints) Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> NSAIDs Member meets <u>ONE</u> of the following: Member tried and failed, has a contraindication, or intolerance to <u>ONE</u> of the following: Cimzia[®] Other Tumor Necrosis Factor (TNF) blocker medication approved for treatment of Active Non-

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Diagnosis: Active Polyarticular Juvenile Idiopathic Arthritis					
Dosing: Oral: Rinvoq® or Rinvoq® LQ					
Patient Age	Patient Weight	Rinvoq® LQ	Rinvoq®		
	10 kg to < 20 kg	3 mg (3 mL) twice daily	Not Recommended		
2 to < 18 years of ag	ge 20 kg to \leq 30 kg	4 mg (4 mL) twice daily	Not Recommended		
	> 30 kg	6 mg (6 mL) twice daily	15 mg once daily		
≥ 18 years of age	N/A	6 mg (6 mL) twice daily	15 mg once daily		
☐ Member has a	a diagnosis of polyarticular j	uvenile idiopathic arthritis			
Prescribed by					
☐ Member is 2					
☐ Member has t					
months cyclosporine					
□ hydroxycl	1				
	leflunomide				
 methotrexate Non-steroidal anti-inflammatory drugs (NSAIDs) oral corticosteroids 					
□ sulfasalaz					
tacrolimu	5				
☐ Member mee	ts ONE of the following:				
Member t	ried and failed, has a contrai	indication, or intolerance to ON	E of the following:		
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					
				auaiii □ Enbre	_
		T) 1-1	1 for the star and of A ations		
Polya	rticular Juvenile Idiopathic A				
☐ Member has been established on Rinvoq® for at least 90 day					
	the past 130 days (verified by				
chart notes or pharmacy paid claims)					

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.