## SENTARA COMMUNITY PLAN (MEDICAID)

## 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 1-800-750-9692</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>

<b><u>Drug Requested</u></b> : select one drug below		
□ Rinvoq® (upadacitinib)	□ Rinvoq® LQ (upadacitinib)	
MEMBER & PRESCRIBER INFORMAT	ΓΙΟΝ: Authorization may be delayed if incomplete.	
Member Name:	_	
Member Sentara #:	Date of Birth:	
Prescriber Name:		
Prescriber Signature:		
Office Contact Name:		
Phone Number:	Fax Number:	
NPI #:		
DRUG INFORMATION: Authorization may		
Drug Name/Form/Strength:		
Dosing Schedule:	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight (if applicable):	Date weight obtained:	
<b>NOTE:</b> The health plan considers the use of concommunomodulator (e.g., Dupixent, Entyvio, Humira, indications to be experimental and investigational. Sa established and will <b>NOT</b> be permitted.		
CLINICAL CRITERIA: Check below all that support each line checked, all documentation, include provided or request may be denied.	apply. All criteria must be met for approval. To ling lab results, diagnostics, and/or chart notes, must be	

□ Diagnosis: Moderate-to-Severe Rheumatoid Arthritis  Dosing: Oral: Rinvoq®: 15 mg once daily						
□ N	☐ Member is at least 18 years of age and have a diagnosis of rheumatoid arthritis (RA)					
	☐ Use in combination with other JAK inhibitors, biologic DMARDS, or potent immunosuppressants such as azathioprine or cyclosporine is not recommended					
□ T	rial and failure of,	contraindication	n, or adver	se reaction to methotrexa	ate	
□ T	☐ Trial and failure of <u>TWO (2)</u> of the preferred drugs below:					
	☐ Humira <sup>®</sup>		□ Enbre	☐ Enbrel®		Infliximab
□ Dia	agnosis: Active	Psoriatic Ar	thritis			
Do	sing: Oral: Rin	nvoq® or Rin	voq® LQ			
P	Patient Age	Patient V	Veight	Rinvoq® LQ		Rinvoq®
		10  kg to < 20  kg	κg	3 mg (3 mL) twice daily	y	Not Recommended
2 to < 1	8 years of age	20  kg to < 30  kg	κg	4 mg (4 mL) twice daily	y	Not Recommended
		> 30 kg		6 mg (6 mL) twice daily	y	15 mg once daily
≥ 18 years of age N/A			N/A		15 mg once daily	
☐ Member has a diagnosis of active psoriatic arthritis						
☐ Member is 2 years of age or older						
Use in combination with other JAK inhibitors, biologic DMARDS, or potent immunosuppressants such as azathioprine or cyclosporine is not recommended						
□ T	☐ Trial and failure of, contraindication, or adverse reaction to methotrexate					
□ T	☐ Trial and failure of <u>TWO (2)</u> of the preferred drugs below:					
	□ Humira <sup>®</sup> □		□ Enbrel <sup>®</sup>			Infliximab
	ngnosis: Moders sing: Oral: Rin		-		nce	daily if inadequate response
N	Member is 12 years	of age or older	weighing a	at least 40kg		

	Prior documented trial and failure of 8 weeks for e	each trial (or contraindication) of:
	☐ One (1) topical corticosteroid of medium to his	gh potency (e.g., mometasone, fluocinolone)
	☐ One (1)) topical calcineurin inhibitors (tacroli	
	☐ Trial and failure of Dupixent®	1
	☐ Trial and failure of Adbry®	
	•	n with other IAV inhibitors higheria DMADDs on
	potent immunomodulators such as azathioprine or	on with other JAK inhibitors, biologic DMARDs, or cyclosporine is not recommended.
	Diagnosis: Moderate-to-Severe Ulcerative	Colitis (UC)
1	<b>Dosing: Oral: Rinvoq</b> <sup>®</sup> : Induction - 45 mg once may increase to 30 mg once daily in patients with refan adequate response is not achieved with the 30 mg maintain response.	ractory, severe, or extensive disease. Discontinue if
	Member is at least 18 years of age and have a diag	mosis of moderate to severe active ulcerative colitis
	Trial and failure of <b>BOTH</b> of the preferred drugs b	pelow:
	☐ Humira <sup>®</sup>	□ Infliximab
	Member is <u>NOT</u> receiving Rinvoq <sup>®</sup> in combination potent immunomodulators such as azathioprine or	on with other JAK inhibitors, biologic DMARDs, or cyclosporine is not recommended
	Diagnosis: Moderate-to-Severe Active Cro	hn's Disease (CD)
]	<b>Dosing: Oral: Rinvoq®:</b> Induction - 45 mg once	e daily for 12 weeks; Maintenance -15 mg once
(	daily; may increase to 30 mg once daily in patients w	with refractory, severe, or extensive disease.
	Discontinue if an adequate response is not achieved	with the 30 mg dose; use the lowest effective dose
	needed to maintain response.  Member is at least 18 years of age and have a diag	mosis of moderate to severe active Crohn's disease
	·	
	·,,,	
	Trial and failure of <b>BOTH</b> of the preferred drugs b	pelow:
	☐ Humira <sup>®</sup>	□ Infliximab
	Member is <u>NOT</u> receiving Rinvoq <sup>®</sup> in combinatio or potent immunomodulators such as azathioprine	
	Diagnosis: Active Ankylosing Spondylitis	
Dosing: Oral: Rinvoq®: 15 mg once daily		
	Member is at least 18 years of age and have a diag	nosis of ankylosing spondylitis

	☐ Trial and failure of <b>TWO (2)</b> of the <b>PREFERRED</b> drugs below:				
	☐ Humira <sup>®</sup>		□ Enbrel®		□ Infliximab
				on with other JAK inle or cyclosporine is no	nibitors, biologic DMARDs, ot recommended
□ Diagnosis: Active Non-Radiographic Axial Spondyloarthritis  Dosing: Oral: Rinvoq®: 15 mg once daily					
	Member is at least	18 years of age a	and have a diag	gnosis of ankylosing	spondylitis
	Trial and failure of	BOTH of the pr	referred drugs	below:	
	☐ Humira <sup>®</sup>			☐ Infliximab	
		•		on with other JAK inlocyclosporine is not in	nibitors, biologic DMARDs, or recommended
u D	iagnosis: Polyar	ticular Juven	ile Idiopath	nic Arthritis	
Dosing: Oral: Rinvoq® or Rinvoq® LQ					
Pati	ent Age	Patient Wei	ght Ri	invoq® LQ	Rinvoq®
		10  kg to < 20  l	xg 3 1	ng (3 mL) twice daily	Not Recommended
,		20 kg to < 30 kg		ng (4 mL) twice daily	Not Recommended
		> 30 kg	6 r	ng (6 mL) twice daily	15 mg once daily
$\geq$ 18 years of age $N/A$		N/A	N/	A	15 mg once daily
	Member has a diagram	nosis of polyarti	cular juvenile	idiopathic arthritis	
			3	1	
_	Member is 2 years	of age or older	J	1	
	-	with other JAK	inhibitors, bio	ologic DMARDS, or	potent immunosuppressants such
	Use in combination as azathioprine or c	with other JAK yclosporine is n	inhibitors, bio	ologic DMARDS, or	
	Use in combination as azathioprine or c	with other JAK yclosporine is n contraindication	inhibitors, bid ot recommend n, or adverse r	ologic DMARDS, or ged eaction to methotrexa	

		Diagnosis: Giant Cell Arteritis Dosing: Oral: Rinvoq®: 15 mg once daily
		Member is 18 years of age or older
		Member has a diagnosis of giant cell arteritis
		Use in combination with other JAK inhibitors, biologic DMARDS, or potent immunosuppressants such as azathioprine or cyclosporine is not recommended
M	ed	ication being provided by Specialty Pharmacy - PropriumRx

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

<sup>\*</sup>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.