## SENTARA COMMUNITY PLAN (MEDICAID)

## MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions:</u> 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; fax to 1-844-305-2331. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization will be delayed.

## **Infliximab Category (MEDICAL)**

**PREFERRED** 

**DRUG REQUESTED:** (Select drug below)

☐ Infliximab (JI745) NDC (57894-0160-01)					
NON-PREFERRED					
□ Avsola <sup>™</sup> (infliximabaxxq) (Q5121)	(infliximab- dyyb)	□ Remicade® (infliximab) (J1745)	□ Renflexis® (infliximab-abda)		
	(Q5103)	NDC (57894-0030-01)	(Q5104)		
MEMBER & PRESCI	RIBER INFORMATIO	ON: Authorization may be o	delayed if incomplete.		
Member Name:					
Member Sentara #:	Member Sentara #: Date of Birth:				
Prescriber Name:					
Prescriber Signature:	rescriber Signature: Date:				
Office Contact Name:					
Phone Number:	Phone Number: Fax Number:				
DEA OR NPI #:					
DRUG INFORMATIO					
Drug Form/Strength:					
Dosing Schedule:	Oosing Schedule: Length of Therapy:				
Diagnosis:	osis: ICD Code, if applicable:		le:		
Weight:	ight: Date:				
☐ Standard Review. In che or the member's ability to		ne does not jeopardize the life and would not subject the m			

- Effective July 1, 2023 per DMAS Infliximab is the preferred infliximab product. Remicade®, Inflectra®, Avsola®, Renflexis® are non-preferred.
- If requesting preferred drug, Infliximab: please check diagnosis and enter the dosing below that applies. No additional prior authorization criteria is required.
- If requesting a non-preferred drug, Renflexis®, Remicade®, Inflectra® or Avsola® please complete all of the required prior authorization criteria.

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DIAGNOSIS	Recommended Dose			
□ Ankylosing Spondylitis (AS) Dosing:	• 5mg/kg at week 0, 2 and 6, then every 6 weeks thereafter			
☐ Crohn's Disease (CD) Dosing:	• 5mg/kg at week 0, 2 and 6, then every 8 weeks thereafter. Max dose 10mg/kg every 8 weeks			
<ul> <li>□ Pediatric Crohn's Disease (CD)</li> <li>□ Age ≥ 6 years</li> <li>□ Dosing:</li></ul>	• 5mg/kg at week 0, 2 and 6 weeks, then every 8 weeks thereafter			
☐ Plaque Psoriasis (Ps) Dosing:	• 5mg/kg at week 0, 2 and 6, then every 8 weeks thereafter			
☐ Psoriatic Arthritis (PsA) Dosing:	• 5mg/kg at week 0, 2 and 6, then every 8 weeks thereafter			
□ Rheumatoid Arthritis (RA) in combination with methotrexate  Dosing:	• 3mg/kg at week 0, 2 and 6, then every 8 weeks thereafter. Max dose 10mg/kg every 8 weeks or 3mg/kg every 4 weeks			
☐ Ulcerative Colitis (UC) Dosing:	• 5mg/kg at week 0, 2 and 6, then every 8 weeks thereafter			
<ul> <li>□ Pediatric Ulcerative Colitis</li> <li>□ Age ≥ 6 years</li> <li>Dosing:</li></ul>	• 5mg/kg at week 0, 2 and 6, then every 8weeks thereafter			
CLINICAL CRITERIA: Check below all that apply. All criteria/diagnosis 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. If requesting an increase in dose, recent lab values and symptoms documenting active disease must be submitted with request.				
□ Diagnosis: Rheumatoid Arthritis or Psoriatic Arthritis				
□ Check diagnosis: □ Rheumatoid Arthritis OR  AND	□ Psoriatic Arthritis			

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	Prescriber is a Rheumatologist					
	AND					
	Trial and failure of <b>ONE</b> of the <b>PREFERRED</b> drugs below:					
	□ azathioprine	□ hydroxychloroqu	uine 🗆	6-mercaptopurine	□ methotrexate	
	□ leflunomide	□ aminosalicylates		auranofin	□ sulfasalazine	
	□ Other					
	AND					
	Trial and failure to Humira®	or Enbrel® AND Inf	liximab the	erapy		
o I	Diagnosis: Ankylosing S	Spondylitis				
	Diagnosed for active ankyle	osing spondylitis				
	AND					
	Prescribed by or in consulta	tion with a Rheumat	ologist			
	AND					
	Trial and failure, contraindication, or intolerance to <u>TWO</u> NSAIDs					
	<u>AND</u>					
	Trial and failure of <b>ONE</b> of the <b>PREFERRED</b> drugs below:					
	□ Humira <sup>®</sup> □ Enbrel <sup>®</sup>					
	AND					
	Trial and failure to Infliximab therapy					
o I	□ Diagnosis: Plaque Psoriasis					
	Diagnosed for Plaque Psoriasis					
	AND					
	Prescribed by or in consultation with a Dermatologist					
	AND					
	Trial and failure of <b>ONE</b> of the <b>PREFERRED</b> drugs below:					
	□ acitretin	□ cyclosporii	ne	□ methotrex	ate	
	AND					

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(	_	Trial and failure to Humira® or Enbrel® AND Infliximab therapy				
<b>3</b>	Diagnosis: Crohn's Disease OR Ocular Sarcoidosis - moderate to severe with inadequate response to:					
[	_	Diagnosed for:				
		□ Crohn's Disease	OR □ O	cular Sarcoidosis		
		AND				
[		Prescribed by or in consultat	ion with a <b>Gastroenterol</b>	ogist		
		<u>OR</u>				
[		Prescribed by or in consultat	ion with an <b>Ophthalmol</b> o	ogist		
		<u>AND</u>				
(	_	Inadequate response to high	dose steroids (e.g.,40-60 a	ng prednisone)		
		<u>AND</u>				
[		Trial and failure of <b>ONE</b> of	the <b>PREFERRED</b> drugs	below:		_
		□ hydroxychloroquine	□ 6-mercaptopurine	□ methotrexate		azathioprine
		□ aminosalicylates	□ auranofin	□ sulfasalazine		leflunomide
		Other:				
		AND				
[	_	Trial and failure to Humira®	AND Infliximab therapy	for Crohn's disease in	dica	tion
	_	D'		1'4' 1'		
	□ Diagnosis: Moderate-to-severe Ulcerative Colitis disease					
[		Diagnosed for moderate-to-severe Ulcerative Colitis				
	AND					
[	☐ Prescribed by or in consultation with a <b>Gastroenterologist</b>					
	<u>AND</u>					
(	<b></b>	Inadequate response to high	dose steroids (e.g.40-60 n	ng prednisone)		
	<u>AND</u>					
	(Continued on next page)					

	Trial and failure of <b>ONE</b> of	the <u>PREFERRED</u> drugs	below:	
	□ hydroxychloroquine	□ 6-mercaptopurine	□ methotrexate	□ azathioprine
	□ aminosalicylates	□ auranofin	□ sulfasalazine	□ leflunomide
	□ Other:			
	AND			
	Γrial and failure to Humira <sup>®</sup>	AND Infliximab therapy		
Me	dication being provide	d by (check below that a	applies):	
	ocation/site of drug admini	stration:		
NI	PI or DEA # of administeri	ng location:		
	<u>OR</u>			
□ Sp	ecialty Pharmacy - Propri	umRx		
standard s a lack egain n		nember to adverse health ously jeopardize the life o	consequences. Sentar or health of the memb	a Health's definition of urgent er or the member's ability to
	ese of samples to initial	e therapy aves not mo	eei siep euii/ preui	unorization Crueria.
* <u>Prev</u>	ious therapies will be ve	erified through pharn	nacy paid claims o	or submitted chart notes.*