## 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 (including phone and fax #s) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

## **Infliximab Category (PHARMACY)**

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

☐ Infliximab NI  NON-PRE  ☐ Inflectra® (infliximab- dyyb)	OC (57894-0160-01) FERRED				
□ Inflectra®	FERRED				
*requires authorization under medical benefit	□ Remicade® (infliximab) NDC (57894-0030-01)	□ Renflexis® (infliximab-abda)			
BER INFORMATION	<b>ON:</b> Authorization may be	delayed if incomplete.			
	Date of B	irth:			
		Date:			
	Fax Number:				
Dosing Schedule:		Length of Therapy:			
Diagnosis:		ICD Code, if applicable:			
	Date:				
	Authorization may be ng this box, the timefram	Fax Number: Fax Number:			

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- 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.

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
□ Pediatric Crohn's Disease (CD)  Age ≥ 6 years  Dosing:	• 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.

Has the member been approved for Infliximab, Remicade or Renflexis previously	y thro	ough	the S	Senta	ıra
medical department?		Yes		l N	o

□ I	□ Diagnosis: Rheumatoid Arthritis or Psoriatic Arthritis						
	Check diagnosis:						
	□ Rheumatoid Arthritis	OR	□ F	Psoriatic Arthritis			
	AND						
	Prescriber is a <b>Rheumatole</b>	ogist					
	AND						
	Trial and failure of <b>ONE</b> o	f the <u>PREFERRED</u> o	drugs belov	V:			
	□ azathioprine	□ hydroxychloroqu	uine 🗖	6-mercaptopurine	□ methotrexate		
	□ leflunomide	<ul><li>aminosalicylates</li></ul>	;	auranofin	□ sulfasalazine		
	□ Other:						
	AND						
	Trial and failure to Humira	® or Enbrel® <b>AND</b> In	fliximab th	erapy			
u I	Diagnosis: Ankylosing	Spondylitis					
	Diagnosed for active anky	losing spondylitis					
	<u>AND</u>						
	Prescribed by or in consult	ation with a Rheuma	tologist				
	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							
□ Diagnosis: Plaque Psoriasis							
	☐ Diagnosed for Plaque Psoriasis						
	<u>AND</u>						
	☐ Prescribed by or in consultation with a Dermatologist						
	AND						

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☐ Trial and failure of <u>ONE</u> of the <u>PREFERRED</u> drugs below:						
	□ acitretin	□ cyclosporine		□ methotrexate		
	AND					
	Trial and failure to Humira®	or Enbrel <sup>®</sup> <b>AND</b> Inflixi	mab therapy			
	Diagnosis: Crohn's Dise severe with inadequate i		·coidosis - mo	derate to		
	Diagnosed for:					
	□ Crohn's Disease	OR 🗅	Ocular Sarcoid	osis		
	AND		1			
Ц	Prescribed by or in consultati	ion with a Gastroenter	ologist			
	OR  Prescribed by or in consultati	ion with an <b>Onhthalm</b> e	alogist			
_	AND	on with an <b>Opithani</b> lo	nogist			
	Inadequate response to high	dose steroids (e.g.,40-60	) mg prednisone	)		
	AND	(	·8 [	,		
	Trial and failure of <b>ONE</b> of t	he <u>PREFERRED</u> drug	gs below:			
	□ hydroxychloroquine	□ 6-mercaptopurine	□ methotrexa	te azathioprine		
	□ aminosalicylates	□ auranofin	u sulfasalazir	e leflunomide		
	Other:					
	AND					
	☐ Trial and failure to Humira® <b>AND</b> Infliximab therapy for Crohn's disease indication					
□ 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>						
	☐ Inadequate response to high dose steroids (e.g.40-60 mg prednisone)					
	AND					

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	☐ Trial and failure of <u>ONE</u> of the <u>PREFERRED</u> drugs below:						
	□ hydroxychloroquine	□ 6-mercaptopurine	□ methotrexate	□ azathioprine			
	□ aminosalicylates	□ auranofin	□ sulfasalazine	□ leflunomide			
	Other:						
0	AND  ☐ Trial and failure to Humira® AND Infliximab therapy						
M	edication being provide	d by (check below that	t applies):				
	Location/site of drug administration:						
	NPI or DEA # of administering location:						
	<u>OR</u>						
	Specialty Pharmacy - Propri	umRx					
a star lack	~	he member to adverse he	ealth consequences. Ser	on Department if they believe ntara's definition of urgent is a or the member's ability to			

\*\*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. \*