

INFLIXIMAB Federal Employee Program. PRIOR APPROVAL REQUEST

Send completed form to: Service Benefit Plan **Prior Approval** P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 **Attn. Clinical Services**

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

physician portion and submit this completed form.		, , , , , ,	3	, ,	Fax: 1-877-37	<u>′8-472</u>	
Patient Inform	Provider Information (required)						
Date:			Provider Name:				
Patient Name:			Specialty:		NPI:		
Date of Birth:	Sex: □Male	□Female	Office Phone:		Office Fax:		
Street Address:			Office Street Address:				
City:	State:	Zip:	City:	Sta	ate: Zip:		
Patient ID: R			Physician Signature:				
IX	P:	HYSICIAN	COMPLETES				
Please select medication:	NOTE: Form m	ust be complet	ed in its entirety for proc	essing			
☐Avsola (infliximab-axxq) ☐Infliximab☐Inflectra (infliximab-dyyb) ☐Remicade (infliximab-dyyb)			□Renflexis (infliximab-abda)				
**Check www.fepblue.org/formulary to	confirm which medica	ation is part of th	e patient's benefit				
months for ALL other diagnod □YES - this is a PA renewal for □NO - this is INITIATION of the state of the st	or CONTINUATI of therapy, please and eric? Brand rior to initiating the ve an active or later are patient started trace infections? Ye is B virus (HBV) in as (HBV) infection accines while on the b-axxq) or Renflex	ION of therapy nswer the quest IGeneric erapy?	y, please answer the questions below: No infection?	□Later c of this m y started tr ave an into	nt TB*	No	
9. Will the requested medication be *If YES, please specify the 1 *DMARDs: Actemra, Avsolution Olumiant, Orencia, Otezla, I Sotyktu, Stelara, Taltz, Trem	nedication: , Cimzia, Cosentyx, Remicade, Renflexis,	Enbrel, Entyvio , Riabni, Rinvoq	o, Humira or a Humira bios	imilar, Ilu	mya, Inflectra, Kevzara, K		
10. What is the patient's weight in	n either pounds or l	kilograms?	lbs <u>OR</u>		kg		
11. Which dosing regimen is the							
12. What is the patient's diagnosi		v <u>x</u>					
☐Behcet's syndrome ☐Pyoderma gangrenosum ☐Ankylosing spondylitis (AS a. Is the patient's condition	□Sarcoidosis □Takayasu's art S) / axial spondyloa	teritis 🗖 G1	idradenitis suppurativa (H ranulomatosis w/polyangi		ener's granulomatosis)		
			vo non-steroidal anti-infla ed anti-inflammatory dose			4 wee	

PLEASE PROCEED TO PAGE 2 FOR ADDITIONAL DIAGNOSES

PAGE 1 of 3



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PAGE 2 – PHYSICIAN COMPLETES						
Patient Name: DOB: Patient ID: R						
□Crohn's disease (CD)						
a. Does the patient have moderate to severely active Crohn's disease (CD)? ☐Yes ☐No						
b. Age 6-17 : Will the patient be current on all vaccinations prior to initiating therapy? □Yes □No						
c. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to convention therapy for Crohn's disease (CD)? Yes No						
□Juvenile idiopathic arthritis (JIA)						
a. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to at least a month trial of a self-injectable TNF inhibitor for juvenile idiopathic arthritis (JIA)? □Yes □No						
□Plaque psoriasis (PsO)						
a. Does the patient have severe plaque psoriasis that covers at least 5% of body surface area (BSA) or affects crucial body areas such as hands, feet, face, neck, scalp, genitals/groin, and intertriginous areas? No						
b. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to convention systemic therapy? Tyes* (*If YES, please select one of the following below) No						
□Inadequate response □Intolerance or contraindication □Has not tried conventional systemic therapy						
c. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to photother Inadequate response Intolerance or contraindication Has not tried phototherapy						
□Psoriatic arthritis (PsA)						
a. Is the patient's psoriatic arthritis active? □Yes □No						
b. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to a 3-mont trial of at least one conventional DMARD? □Yes □No						
□Rheumatoid arthritis (RA)						
a. Does the patient have moderate to severely active rheumatoid arthritis (RA)? ☐Yes ☐No						
b. Has the patient had an inadequate response to at least a 3 month trial of methotrexate despite adequate dosing (i.e., titra 20mg/week)? □Yes □No*						
* $If NO$, does the patient have a contraindication or intolerance to methotrexate? \square Yes \square No						
c. Does the patient have a contraindication or intolerance to leflunomide? ☐Yes ☐No*						
* $If NO$, will the patient receive concurrent therapy with either methotrexate or leflunomide? \square Yes \square No						
□Ulcerative colitis (UC)						
a. Does the patient have moderate to severely active ulcerative colitis (UC)? ☐Yes ☐No						
b. Age 6-17 : Will the patient be current on all vaccinations prior to initiating therapy? □Yes □No						
c. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to convention therapy for ulcerative colitis (UC)?						
□Uveitis						
a. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to a trial of immunosuppressive therapy for uveitis? ☐Yes ☐No						
Other diagnosis (nlease specify):						

PAGE 2 of 3



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physician portion and submit this completed form.	Fax: 1-877-378-4727								
Patient Information (required)			Provider Information (required)						
Date:			Provider Name:						
Patient Name:			Specialty:		NPI:				
Date of Birth:	Sex: □Male	□Female	Office Phone:	Office Phone: Office Fax:					
Street Address:			Office Street Address:	:					
City:	State:	Zip:	City:	Sta	te:	Zip:			
Patient ID: R	1 1 1		Physician Signature:						
PHYSICIAN COMPLETES									

CONTINUATION OF THERAPY (PA RENEWAL) NOTE: Form must be completed in its **entirety** for processing Please select medication: □Avsola (infliximab-axxq) □Infliximab □Renflexis (infliximab-abda) □Inflectra (infliximab-dyyb) □Remicade (infliximab) **Check www.fepblue.org/formulary to confirm which medication is part of the patient's benefit 1. Has the patient been on the requested medication continuously for the last 4 months for Rheumatoid Arthritis OR for the last 3 months for ALL other diagnoses excluding samples? Please select answer below: □NO – this is INITIATION of therapy, please answer the questions on PAGE 1 **TYES** – this is a PA renewal for **CONTINUATION** of therapy, please answer the questions below: 2. Is this request for brand or generic? □Brand □Generic 3. What is the patient's diagnosis? □Ankylosing spondylitis (AS) / axial spondyloarthritis ☐Psoriatic arthritis (PsA) ☐Behcet's syndrome □Pyoderma gangrenosum □Crohn's disease (CD) □Rheumatoid arthritis (RA) ☐ Hidradenitis suppurativa (HS) □ Sarcoidosis ☐Granulomatosis w/polyangiitis (Wegener's granulomatosis) ☐Takayasu's arteritis ☐ Juvenile idiopathic arthritis (JIA) □Ulcerative colitis (UC) ☐Plaque psoriasis (PsO) **□**Uveitis □Other (*please specify*): _____ 4. Has the patient's condition improved or stabilized with therapy? \(\sigma Y \)es 5. Does the patient have any active infections including tuberculosis (TB) and hepatitis B virus (HBV)? \(\preceq \text{Yes} \) 6. Will the patient be given live vaccines while on this therapy? □Yes 7. Will the requested medication be used in combination with another biologic *DMARD or targeted synthetic DMARD? □Yes* *If YES, please specify the medication: *DMARDs: Actemra, Avsola, Cimzia, Cosentyx, Enbrel, Entyvio, Humira or a Humira biosimilar, Ilumya, Inflectra, Kevzara, Kineret, Olumiant, Orencia, Otezla, Remicade, Renflexis, Riabni, Rinvoq, Rituxan, Ruxience, Siliq, Simponi/Simponi Aria, Skyrizi, Spevigo, Sotyktu, Stelara, Taltz, Tremfya, Truxima, and Xeljanz 8. What is the patient's weight in either pounds or kilograms? 9. Which dosing regimen is the patient on (specify dose and frequency)? ___

PAGE 3 of 3

10. Did the patient have an inadequate treatment response to the initial dosing regimen, and is therefore considered to be a non-