

INFLIXIMAB PRIOR APPROVAL REQUEST

Send completed form to: FAX: 855-895-3504 FOR URGENT FAX: 844-244-0226

Federal Employee Program PRIOR APPROVAL REQUEST

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.

Patient Information (required)				Provider Information (required)				
Date:				Provider Nan	ne:			
Patient Name:				Specialty:			NPI:	
Date of Birth:		Sex: Male	Female	Office Phone:			Office Fax:	
Street Address:		-		Office Street	Address:			
City:		State:	Zip:	City: State		State:	z: Zip:	
Patient ID: R	Patient ID:			Physician Signature:				
K		I	PHYSICIAN C	OMPLETE	S			
		NOTE: Form n	nust be completed	d in its entire t	v for processin	10		
		NOTE: TORRE	tust be completed	a iii its chtii c i	y for processin	<u>.e</u>		
Please select me	dication:							
□Avsola (inflix	ximab-axxq)	□Inflectra (infl	iximab-dyyb)	□Inflixin	ıab	□Renfle	exis (inflixi	imab-abda)
**Check www.fepbl	lue.org/formulary to	confirm which medic	cation is part of the	patient's benefit				
☐ YES - this : ☐ NO - this is 2. Is this request 3. Patient 6-17 \(\) 4. Has the patient *If YES, do *If Latent 5. Does the patient *If YES, ha *If	LL other diagnors is a PA renewal for a PA renewal for a INITIATION of a for brand or generated by the state of Age: With the state of Ag	or CONTINUAT of therapy, please a eric? Brand ill the patient be c sis (TB) test prior we an active or late ient started treatm re infections? TY is B (HBV) infect been ruled out or accines while on t	ATON of therapy, answer the questing Generic current on all vacce to initiating there ent TB infection from the infect for the infect for the infect for the infect for the patient all this therapy?	please answer ions below: cinations prior apy?	the questions of the questions of the questions of the property of the questions of the que	nerapy? B* □7 edication HBV infe	□Yes □ Test was ne n? □Yes ection? □Y	□No Yes □No
*If YES, pl *DMARD Orencia, (netic DMARD? [lease specify: 0 includes: Actemra Otezla, Remicade, F and Xeljanz	□Yes* □No a, Avsola, Cimzia, C	Cosentyx, Enbrel, 1	Entyvio, Humir	a, Ilumya, Inflec	ctra, Kevz	zara, Kinere	t, Olumiant,
9. What is the pa	atient's weight in	either pounds or l	cilograms?	lbs	<u>OR</u>		_ kg	
10. What dosing	regimen is the pa	atient on (specify d	ose and frequency)?				
11. What is the p	patient's diagnosis	s?						
☐Behcet's s	syndrome			□Hidrade	nitis suppurativ	va	□ Sarcoid	losis
□Granulom	atosis w/polyangi	iitis (Wegener's g	ranulomatosis)	□Pyodern	na gangrenosur	m	□Takaya	su's arteritis
□Ankylosir	ng Spondylitis (AS	S) / axial spondylo	oarthritis					
a. Is the	patient's condition	on active? Yes	□No					
	he patient had an a period in total at							

PLEASE PROCEED TO PAGE 2 FOR ADDITIONAL DIAGNOSES

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BlueShield. INFLIXIMAB Federal Employee Program. PRIOR APPROVAL REQUEST

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PAGE 2 - PHYSICIAN COMPLETES						
Patient Name:	DOB:	Patient ID: R				
□Crohn's Disease (CD)						
a. Does the patient have moderate t	to severely active Crohn's	s disease? □Yes □No				
b. Does the patient have a contraind therapy for Crohn's disease?	•	either an inadequate response or intolerance to conventional				
☐Juvenile Idiopathic Arthritis (JIA)						
 Does the patient have a contraind month trial of a self-injectable Tl 		either an inadequate response or intolerance to at least a three idiopathic arthritis? No				
□Plaque Psoriasis (Ps)						
 a. Does the patient have severe place areas such as hands, feet, face, no 		at least 5% of body surface area (BSA) or affects crucial body and intertriginous areas? ☐Yes ☐No				
b. Does the patient have a contraindic therapy?		her an inadequate response or intolerance to conventional systemic below) \square No				
		☐ Has not tried conventional systemic therapy				
-	•	either an inadequate response or intolerance to phototherapy?				
☐Inadequate response ☐Intol	erance or contraindication	☐Has not tried phototherapy				
□Psoriatic Arthritis (PsA)						
a. Is the psoriatic arthritis active?						
b. Does the patient have a contraine of at least one conventional disea		either an inadequate response or intolerance to a three month triantic drug (DMARD)?				
□Rheumatoid Arthritis (RA)						
a. Does the patient have moderate t	o severely active rheuma	toid arthritis? □Yes □No				
b. Has the patient had an inadequat titrated to 20mg/week)? □Yes	e response to at least a th	ree-month trial of methotrexate despite adequate dosing (i.e.,				
*If NO, does the patient have	a contraindication or inte	olerance to methotrexate? □Yes □No				
c. Does the patient have a contraince	lication or intolerance to	leflunomide? □Yes □No*				
*If NO, will the patient receive	ve concurrent therapy wit	h either methotrexate or leflunomide? ☐Yes ☐No				
□Ulcerative Colitis (UC)						
a. Does the patient have moderate t	o severely active ulcerati	ve colitis? □Yes □No				
b. Does the patient have a contraind therapy for ulcerative colitis?		either an inadequate response or intolerance to conventional				
□Uveitis						
 a. Does the patient have a contraint immunosuppressive therapy? 		either an inadequate response or intolerance to a trial of				
□Other diagnosis (please specify):						

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Date:			Provider Mame: (required)				
Patient Name:				Specialty: NPI:		NPI:	
Date of Birth:		Sex: Male	Female	Office Phone:		Office Fax:	
Street Address:				Office Street Address:			
City:		State:	Zip:	City:	S	State:	Zip:
Patient ID:				Physician Signature:			
R	l I	P:	HYSICIAN (COMPLETES			
CONTINUATION OF THERAPY (PA RENEWAL)							
	CON			d in its entirety for pro		AL)	
		NOTE. Politi ini	ust be complete	d in its entirety for pro	ocessing .		
Please select med							
□Avsola (inflix		□Inflectra (inflix		□Infliximab		Renflexis (inf	liximab-abda)
□YES - this is a PA renewal for CONTINUATION of therapy, 2. Is this request for brand or generic? □Brand □Generic 3. What is the patient's diagnosis? □Ankylosing Spondylitis (AS) / axial spondyloarthritis □Behcet's syndrome □Crohn's Disease (CD) □Hidradenitis suppurativa □Granulomatosis w/polyangiitis (Wegener's granulomatosis) □Juvenile Idiopathic Arthritis (JIA) □Plaque Psoriasis (Ps) □Other diagnosis (please specify):				□Psoriatic Arthritis (PsA) □Pyoderma gangrenosum □Rheumatoid Arthritis (RA) □Sarcoidosis □Takayasu's arteritis □Ulcerative Colitis (UC) □Uveitis			
-	•	roved or stabilized			/HDMA 6	-	r
•	· ·		•	s (TB) and hepatitis B	(HBV)? L No	∃Yes □N	0
7. Will the requestargeted synthes *If YES, ple *DMARD Orencia, O	sted medication betic DMARD? case specify: includes: Actemra	JYes* □No , Avsola, Cimzia, Co	ntion with anoth	er biologic *disease-m Entyvio, Humira, Ilumya Ruxience, Siliq, Simponia	odifying a	, Kevzara, Kin	neret, Olumiant,
8. What is the pa	tient's weight in e	either pounds or ki	ilograms?	lbs <u>OR</u>		kg	
9. What dosing re	egimen is the pati	ent on (specify dose	e and frequency)	?			
10. Did the patier responder?		uate treatment res	ponse to the ini	tial dosing regimen, an	d is theref	fore consider	ed a non-

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The information provided on this form will be used to determine the provision of healthcare benefits under a U.S. federal government program, and any falsification of records may subject the provider to prosecution, either civilly or criminally, under the False Claim Acts, the False Statements Act, the mail or wire fraud statutes, or other federal or state laws prohibiting such falsification. **Prescriber Certification:** Lectify all information provided on this form to be true and correct to the best of my knowledge and belief. I understand that the insurer may request a medical record if the information provided herein is not sufficient to make a benefit determination or requires clarification and I agree to provide any such information to the insurer. Inflixmab – FEP MD Fax Form Revised 1/7/2022